=> d his '

(FILE 'HOME' ENTERED AT 14:00:41 ON 01 SEP 2005)

FILE 'HCAPLUS' ENTERED AT 14:01:12 ON 01 SEP 2005 1 US2004253289/PN OR US2004-711162#/AP,PRN

FILE 'REGISTRY' ENTERED AT 14:01:49 ON 01 SEP 2005

FILE 'HCAPLUS' ENTERED AT 14:01:49 ON 01 SEP 2005
L2 TRA L1 1- RN : 4 TERMS

FILE 'REGISTRY' ENTERED AT 14:01:50 ON 01 SEP 2005 L3 4 SEA L2

FILE 'WPIX' ENTERED AT 14:01:54 ON 01 SEP 2005 L4 1 L1

FILE 'HCAPLUS' ENTERED AT 14:02:18 ON 01 SEP 2005

FILE 'REGISTRY' ENTERED AT 14:02:23 ON 01 SEP 2005

=> b hcap;d all l1 FILE 'HCAPLUS' ENTERED AT 14:03:03 ON 01 SEP 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 1 Sep 2005 VOL 143 ISS 10 FILE LAST UPDATED: 31 Aug 2005 (20050831/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

```
L1 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2005 ACS on STN
```

AN 2004:1080523 HCAPLUS

DN 142:16788

ED Entered STN: 17 Dec 2004

TI Natural plant compound with anti-hiv activity

IN Khripach, Vladimir; Altsivanovich, Konstantin; Zabinskii, Vladimir; Samusevich, Mikhail

PA Mikonik Technologies, Ltd., Belarus; Drebsk Comptech, Inc.

SO U.S. Pat. Appl. Publ., 5 pp. CODEN: USXXCO

DT Patent

LA English

IC ICM A61K031-415
ICS A01N043-52; A61K047-00; A61K035-78; A61K009-20; A61K009-48; A61K009-14

INCL 424422000; 424464000; 424465000; 424439000; 424451000; 424489000; 424725000

```
CC
    1-5 (Pharmacology)
    Section cross-reference(s): 11, 17
FAN. CNT 1
                                       APPLICATION NO. DATE
    PATENT NO.
                      KIND
                               DATE
     -----
                               20041216 US 2004-711162 20040828 <--
                      A1
    US 2004253289
ΡI
PRAI US 2004-711162
                               20040828 <--
CLASS
PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
US 2004253289 ICM
                       A61K031-415
                       A01N043-52; A61K047-00; A61K035-78; A61K009-20;
                ICS
                       A61K009-48; A61K009-14
                INCL 424422000; 424464000; 424465000; 424439000; 424451000;
                       424489000; 424725000
                      424/422.000
US 2004253289
                NCL
                ECLA A23L001/30; A61K031/415; A61K031/415+M; A61K045/06 <--
    The invention comprises a method for treatment of HIV-infection and
AB
    related conditions, particularly AIDS, using plant hormone
    24-epibrassinolide, anti-HIV efficacy of which is disclosed.
    epibrassinolide natural plant hormone HIV antiHIV
ST
    Hormones, plant
IT
    RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
    study); OCCU (Occurrence); USES (Uses)
        (brassinosteroids; natural plant compound, 24-epibrassinolide with
       anti-hiv activity)
    Drug delivery systems
IT
        (capsules; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Drug delivery systems
IT
        (coating; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Contraceptives
IT
        (condoms; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Drug delivery systems
IT
        (emulsions, aqueous; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    AIDS (disease)
IT
    Anti-AIDS agents
    Combination chemotherapy
    Drug delivery systems
    Food
    Human
    Human immunodeficiency virus
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
    Natural products, pharmaceutical
IT
    RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
    study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
    Drug delivery systems
IT
        (ointments, creams; natural plant compound, 24-epibrassinolide with
       anti-hiv activity)
    Drug delivery systems
IT
        (powders; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Drug delivery systems
IT
        (solns.; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
IT
    Diet
        (supplements; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Drug delivery systems
IT
        (suppositories, vaginal; natural plant compound, 24-epibrassinolide with
```

```
anti-hiv activity)
    Drug delivery systems
IT
        (suspensions; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
    Drug delivery systems
IT
        (tablets; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
IT
    Vagina
        (tract, protection by HIV-inhibiting 24-epibrassinolid-containing composition;
        natural plant compound, 24-epibrassinolide with anti-hiv activity)
     9068-38-6
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HIV, inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     144114-21-6, HIV protease
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     78821-43-9, 24-Epibrassinolide
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
     study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
     52350-85-3, HIV integrase
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (of HIV, inhibitor; natural plant compound, 24-epibrassinolide with
        anti-hiv activity)
=> b reg;d ide l3 tot
FILE 'REGISTRY' ENTERED AT 14:03:10 ON 01 SEP 2005
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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STRUCTURE FILE UPDATES: 31 AUG 2005 HIGHEST RN 862246-83-1
DICTIONARY FILE UPDATES: 31 AUG 2005 HIGHEST RN 862246-83-1
New CAS Information Use Policies, enter HELP USAGETERMS for details.
```

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

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Structure search iteration limits have been increased. See HELP SLIMITS for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

```
ANSWER 1 OF 4 REGISTRY COPYRIGHT 2005 ACS on STN
L3
RN
     144114-21-6 REGISTRY
   Entered STN: 23 Oct 1992
     Retropepsin (9CI) (CA INDEX NAME)
OTHER NAMES:
    Avian leukosis virus proteinase
     E.C. 3.4.23.16
CN
     Endogenous retroviral proteinase
CN
     FIV proteinase
CN
CN
     Gag Protease
     HIV aspartyl protease
CN
CN
     HIV protease
     HIV proteinase
CN
     HIV-1 aspartyl protease
CN
    HIV-1 aspartyl proteinase
CN
    HIV-1 protease
CN
     HIV-1 proteinase
CN
     HIV-1 virus aspartyl proteinase
CN
    HIV-1 virus protease
CN
     HIV-2 protease
CN
CN
     HTLV proteinase
     HTLV-1 proteinase
CN
     HTLV-I protease
CN
     Human immunodeficiency virus protease
CN
     Mason-Pfizer monkey virus protease
CN
     Moloney murine leukemia virus protease
CN
     Retroproteinase
CN
     Rous sarcoma virus protease
CN
     RSV proteinase
CN
     Simian immunodeficiency virus aspartyl proteinase
CN
     STLV protease
CN
     Unspecified
MF
     COM, MAN
CI
SR
     CA
     STN Files: AGRICOLA, BIOBUSINESS, BIOSIS, CA, CAPLUS, CASREACT, CIN,
LC
       PROMT, TOXCENTER, USPAT2, USPATFULL
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
            4117 REFERENCES IN FILE CA (1907 TO DATE)
             119 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
            4144 REFERENCES IN FILE CAPLUS (1907 TO DATE)
     ANSWER 2 OF 4 REGISTRY COPYRIGHT 2005 ACS on STN
L3
RN
     78821-43-9 REGISTRY
     Entered STN: 16 Nov 1984
ED
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,
     (2\alpha, 3\alpha, 5\alpha, 22R, 23R) -
OTHER NAMES:
     24(R)-Epibrassinolide
CN
     24-epi-Brassinolide
CN
     24-Epibrassinolide
CN
     24-epibrassinolide
CN
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-(2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl)hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     [1R-[1\alpha(1S*,2R*,3R*,4R*),3a\beta,3b\alpha,6a\beta,8\beta,9\beta]
     ,10a\alpha,10b\beta,12a\alpha]]-
CN
     B 1105
     BP 55
CN
```

```
CN Epibrassinolide
```

CN Epibrassinolide R

CN Epin

FS STEREOSEARCH

DR 126721-49-1

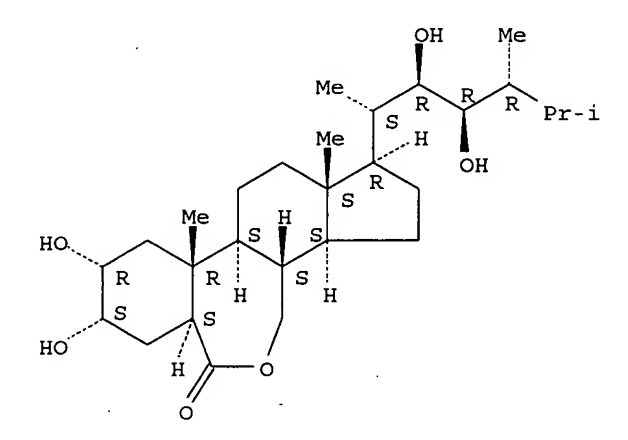
MF C28 H48 O6

CI COM

LC STN Files: AGRICOLA, BEILSTEIN\*, BIOBUSINESS, BIOSIS, CA, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CSCHEM, PROMT, TOXCENTER, USPAT2, USPATFULL

(\*File contains numerically searchable property data)

## Absolute stereochemistry.



## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

315 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

315 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 3 OF 4 REGISTRY COPYRIGHT 2005 ACS on STN

RN 52350-85-3 REGISTRY

ED Entered STN: 16 Nov 1984

CN Integrase (9CI) (CA INDEX NAME)

OTHER NAMES:

CN DNA integrase

CN Enzymes, DNA-recombining, gene int

CN FimE integrase

CN Gene int proteins

CN HIV integrase

CN Proteins, gene int

DR 71850-92-5

MF Unspecified

CI MAN

LC STN Files: AGRICOLA, BIOBUSINESS, BIOSIS, CA, CAPLUS, CEN, CIN, PROMT, TOXCENTER, USPATZ, USPATFULL

## \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

2354 REFERENCES IN FILE CA (1907 TO DATE)

34 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

2364 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 4 OF 4 REGISTRY COPYRIGHT 2005 ACS on STN

RN 9068-38-6 REGISTRY

```
ED
     Entered STN: 16 Nov 1984
     Nucleotidyltransferase, deoxyribonucleate, RNA-dependent (9CI) (CA INDEX
CN
     NAME)
OTHER NAMES:
     Cyscribe reverse transcriptase
CN
     Cyscript
     Reverse transcriptase
CN
     Revertase
CN
CN
     RNA revertase
     RNA-dependent deoxyribonucleate nucleotidyltransferase
CN
     RNA-dependent DNA polymerase
CN
     RNA-directed DNA polymerase
CN
     RNA-instructed DNA polymerase
CN
     SuperScript
CN
CN
     SuperScript II
     ThermoScript
CN
     ThermoScript II
CN
     Unspecified
MF
CI
     MAN
LC
     STN Files:
                  ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO,
       CA, CABA, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST, CIN, CSCHEM,
       EMBASE, IFICDB, IFIPAT, IFIUDB, MSDS-OHS, NAPRALERT, PIRA, PROMT,
       TOXCENTER, USPAT2, USPATFULL
     Other Sources:
                      EINECS**
          (**Enter CHEMLIST File for up-to-date regulatory information)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
            9469 REFERENCES IN FILE CA (1907 TO DATE)
             135 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
            9494 REFERENCES IN FILE CAPLUS (1907 TO DATE)
=> b wpix;d all 14 tot
FILE 'WPIX' ENTERED AT 14:03:17 ON 01 SEP 2005
COPYRIGHT (C) 2005 THE THOMSON CORPORATION
                                              <20050826/UP>
FILE LAST UPDATED:
                            26 AUG 2005
MOST RECENT DERWENT UPDATE:
                                 200555
                                              <200555/DW>
DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE
>>> FOR A COPY OF THE DERWENT WORLD PATENTS INDEX STN USER GUIDE,
    PLEASE VISIT:
 http://www.stn-international.de/training_center/patents/stn_guide.pdf <<<
>>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE
    http://thomsonderwent.com/coverage/latestupdates/
                                                                 <<<
>>> FOR INFORMATION ON ALL DERWENT WORLD PATENTS INDEX USER
    GUIDES, PLEASE VISIT:
    http://thomsonderwent.com/support/userguides/
                                                                 <<<
>>> NEW! FAST-ALERTING ACCESS TO NEWLY-PUBLISHED PATENT
    DOCUMENTATION NOW AVAILABLE IN DERWENT WORLD PATENTS INDEX
    FIRST VIEW - FILE WPIFV.
    FOR FURTHER DETAILS: http://www.thomsonderwent.com/dwpifv <<<
>>> THE CPI AND EPI MANUAL CODES HAVE BEEN REVISED FROM UPDATE 200501.
    PLEASE CHECK:
http://thomsonderwent.com/support/dwpiref/reftools/classification/code-revision/
    FOR DETAILS. <<<
'BIX BI, ABEX' IS DEFAULT SEARCH FIELD FOR 'WPIX' FILE
```

```
ANSWER 1 OF 1 WPIX COPYRIGHT 2005 THE THOMSON CORP on STN
L4
AN
     2005-030192 [03]
                        WPIX
DNC C2005-009587
     Use of 24-epibrassinolide, which is plant hormone belonging to
TI
     brassinosteroid series for the treatment of human immunodeficiency virus
     infection and related diseases.
     B04 D13
DC
     ALTSIVANOVICH, K; KHRIPACH, V; SAMUSEVICH, M; ZABINSKII, V
IN
     (DREB-N) DREBSK COMPTECH INC; (MIKO-N) MIKONIK TECHNOLOGIES LTD
PA
CYC 1
ΡI
     US 2004253289
                     A1 20041216 (200503)*
                                                 5
                                                      A61K031-415
ADT US 2004253289 A1 US 2004-711162 20040828
PRAI US 2004-711162
                          20040828
IC
     ICM A61K031-415
     ICS A01N043-52; A61K009-14; A61K009-20; A61K009-48; A61K035-78;
          A61K047-00
AB
     US2004253289 A UPAB: 20050112
     NOVELTY - Inhibition or treatment of Human Immunodeficiency Virus (HIV)
     infection involves administration of 24-epibrassinolide (EBI) which is a
     plant hormone belonging to brassinosteroid series.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the
     following:
          (1) a pharmaceutical composition comprising 24-epibrassinolide
     optionally in combination with other anti-HIV agents; and
          (2) a food supplement containing 24-epibrassinolide.
          ACTIVITY - Anti-HIV. The efficacy of 24-epibrassinolide (Ia) to
     protect cells against HIV was evaluated in suspensional T-lymphoblastoid
     cell line (MT-4) by Formazan assay based on metabolic reduction of
     3-(4,5-diemthylthiazol-2-yl)-2,5-diphenyltetrazolium bromide. The cells
     infected with HIV-1 (Strain zmb) were incubated with (Ia) (10 - 1 ng/ml).
     By colorimetric analysis, it was found that (Ia) protected the cells
     against HIV-1-cytopathic action.
          MECHANISM OF ACTION - Viral replication inhibitor.
          USE - For the prophylaxis or therapy of AIDS and related diseases;
     and in food supplements (claimed).
          ADVANTAGE - The 24-epibrassinolide is safe and natural steroidal
     plant growth hormone for therapeutic use; exhibits excellent antiviral
     activity, non-toxicity and other positive effects such as blood
     cholesterol lowering activity; reduces cyto-killing properties of viruses;
     increases cell's resistance to the HIV influence; and is a potent
     inhibitor of viral replication.
     Dwq.0/0
FS
     CPI
FA
     AB; DCN
     CPI: B04-J02; B14-A02B1; B14-D03; B14-G01B; D03-H01T2
MC
=> b home
FILE 'HOME' ENTERED AT 14:03:24 ON 01 SEP 2005
=>
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=> b reg;d ide can 17 tot;d que sta 110

FILE 'REGISTRY' ENTERED AT 15:41:12 ON 01 SEP 2005

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TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

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Structure search iteration limits have been increased. See HELP SLIMITS for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

- L7 ANSWER 1 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN
- RN 551928-73-5 REGISTRY
- ED Entered STN: 21 Jul 2003
- CN 5H-Benz[b] indeno[5,4-d] oxepin-5-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,5-dimethyl-4-(methyl-d3)hexyl-6,6,6-d3]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)
- FS STEREOSEARCH
- MF C28 H42 D6 O6
- SR CA
- LC STN Files: CA, CAPLUS, CASREACT

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 139:69422

L7 ANSWER 2 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 461670-12-2 REGISTRY

ED Entered STN: 16 Oct 2002

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl-6,6,6-d3]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8R,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C28 H45 D3 O6

SR CA

LC STN Files: CA, CAPLUS, CASREACT

Absolute stereochemistry.

2 REFERENCES IN FILE CA (1907 TO DATE)

2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 141:221056

REFERENCE 2: 137:263228

L7 ANSWER 3 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 273753-11-0 REGISTRY

ED Entered STN: 29 Jun 2000

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4-

dimethylheptyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS, CASREACT

Absolute stereochemistry.

### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 140:352046

REFERENCE 2: 133:30861

L7 ANSWER 4 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 273753-05-2 REGISTRY

ED Entered STN: 29 Jun 2000

SH-Benz[b] indeno[5,4-d] oxepin-5-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4-dimethylheptyl] hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
(1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS, CASREACT

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 133:30861

L7 ANSWER 5 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 267221-93-2 REGISTRY

ED Entèred STN: 30 May 2000

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF · C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

# \* \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 132:322030

L7 ANSWER 6 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 259104-16-0 REGISTRY

ED Entered STN: 13 Mar 2000

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aR,8R,9S,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 2,3,5-Tri-epi-brassinolide

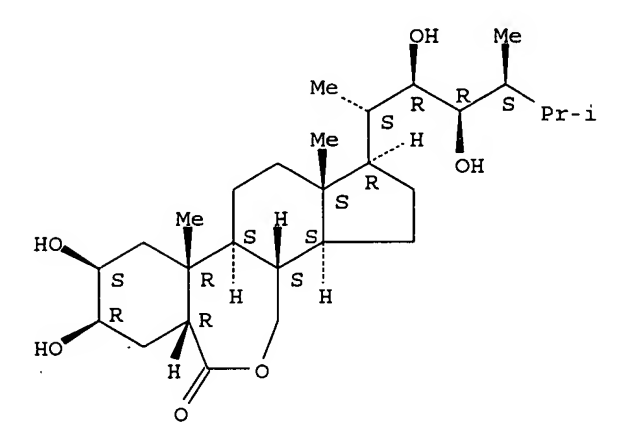
FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.



## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 132:166386

L7 ANSWER 7 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 220401-55-8 REGISTRY

ED Entered STN: 11 Mar 1999

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8R,9S,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 2,3-Di-epi-brassinolide

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS

3 REFERENCES IN FILE CA (1907 TO DATE)

3 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 132:308542

REFERENCE 2: 132:166386

REFERENCE 3: 130:168538

L7 ANSWER 8 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 220401-52-5 REGISTRY

ED Entered STN: 11 Mar 1999

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-

trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9S,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 2-epi-Brassinolide

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS

2 REFERENCES IN FILE CA (1907 TO DATE)

2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 132:308542

REFERENCE 2: 130:168538

L7 ANSWER 9 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 218623-69-9 REGISTRY

ED Entered STN: 29 Jan 1999

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aR,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 5-epi-Brassinolide

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 130:81696

L7 ANSWER 10 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 215502-64-0 REGISTRY

ED Entered STN: 13 Dec 1998

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4-dimethyl-5-(methyl-d3)hexyl-5,6,6,6-d4]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C28 H41 D7 O6

SR CA

LC STN Files: CA, CAPLUS, CASREACT

2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 132:180768

REFERENCE 2: 129:343625

L7 ANSWER 11 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 163514-19-0 REGISTRY

ED Entered STN: 06 Jun 1995

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,

(1R, 3aS, 3bS, 6aS, 8R, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\beta, 5\alpha, 22R, 23R)$  -

OTHER NAMES:

CN 3,24-Diepibrassinolide

FS STEREOSEARCH

MF C28 H48 O6

SR CZ

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 122:310808

L7 ANSWER 12 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 146205-07-4 REGISTRY

ED Entered STN: 26 Feb 1993

CN 5H-Benz[b] indeno[5,4-d] oxepin-5-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl] hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN B-Homo-6-oxaergostan-7-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha, 3\alpha, 5\alpha, 22R, 23R, 24S)$ -

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT

(\*File contains numerically searchable property data)

Absolute stereochemistry.

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

3 REFERENCES IN FILE CA (1907 TO DATE)

3 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 140:352046

REFERENCE 2: 119:95915

REFERENCE 3: 118:147870

L7 ANSWER 13 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 145430-52-0 REGISTRY

ED Entered STN: 21 Jan 1993

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one-4-t, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-4,6a-t2-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one-5,7a,7a-t3 deriv.

CN B-Homo-7-oxaergostan-6-one-5,7a,7a-t3, 2,3,22,23-tetrahydroxy-, (2α,3α,5α,22R,23R)-

FS STEREOSEARCH

MF C28 H45 O6 T3

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

2 REFERENCES IN FILE CA (1907 TO DATE)

2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 124:50719

REFERENCE 2: 118:59960

L7 ANSWER 14 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 140923-40-6 REGISTRY

ED Entered STN: 01 May 1992

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,

(1R, 3aS, 3bS, 6aS, 8R, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\beta, 5\alpha, 22R, 23R, 24S)$  -

OTHER NAMES:

CN 3-Epibrassinolide

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMINFORMRX

(\*File contains numerically searchable property data)

4 REFERENCES IN FILE CA (1907 TO DATE)

4 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 142:276838

REFERENCE 2: 136:243965

REFERENCE 3: 130:168538

REFERENCE 4: 116:211195

L7 ANSWER 15 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 135559-12-5 REGISTRY

ED Entered STN: 16 Aug 1991

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2S,3R,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,

(1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\alpha, 5\alpha, 22S, 23R) -$ 

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT, CHEMINFORMRX (\*File contains numerically searchable property data)

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 115:92706

L7 ANSWER 16 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 128134-34-9 REGISTRY

ED Entered STN: 13 Jul 1990

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one-7-14C, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one-4-14C deriv.

CN B-Homo-7-oxaergostan-6-one-4-14C, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\alpha, 5\alpha, 22R, 23R)$  -

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS, CASREACT

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

# 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 113:59677

L7 ANSWER 17 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 128097-87-0 REGISTRY

ED Entered STN: 06 Jul 1990

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one-7-14C, 1-[(1S,2S,3S,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one-4-14C deriv.

CN B-Homo-7-oxaergostan-6-one-4-14C, 2,3,22,23-tetrahydroxy-, (2α,3α,5α,22S,23S)-

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: CA, CAPLUS, CASREACT

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 113:59677

L7 ANSWER 18 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 115783-59-0 REGISTRY

ED Entered STN: 13 Aug 1988

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one-4-d, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-4,6a-d2-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one-5,7a,7a-d3 deriv.

CN B-Homo-7-oxaergostan-6-one-5,7a,7a-d3, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22R,23R)$ -

FS STEREOSEARCH

MF C28 H45 D3 O6

SR CA

LC STN Files: CA, CAPLUS

2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 118:59960

REFERENCE 2: 109:89818

L7 ANSWER 19 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 113666-77-6 REGISTRY

ED Entered STN: 02 Apr 1988

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1R,2R,3R,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\alpha, 5\alpha, 20R, 22R, 23R, 24S) -$ 

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT, CHEMINFORMRX (\*File contains numerically searchable property data)

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 108:187052

L7 ANSWER 20 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 110611-54-6 REGISTRY

ED Entered STN: 10 Oct 1987

CN B-Homo-7-oxaergostan-6-one-26,26,26,28,28,28-d6, 2,3,22,23-tetrahydroxy-, (2α,3α,5α,22R,23R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-6-oxaergostan-6-one-26,26,26,28,28,28-d6 deriv.

FS STEREOSEARCH

MF C28 H42 D6 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT

(\*File contains numerically searchable property data)

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 107:154590

L7 ANSWER 21 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 110453-84-4 REGISTRY

ED Entered STN: 27 Sep 1987

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\beta,22S,23S,24S)$ - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

OTHER NAMES:

CN (22S, 23S, 24S) - Epibrassinolide

FS STEREOSEARCH

MF C28 H48 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMINFORMRX (\*File contains numerically searchable property data)

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 107:154592

L7 ANSWER 22 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 107853-67-8 REGISTRY

ED Entered STN: 02 May 1987

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetra(hydroxy-d)-,

 $(2\alpha, 3\alpha, 5\alpha, 22R, 23R)$  - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

FS STEREOSEARCH

MF C28 H44 D4 O6

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 106:172256

L7 ANSWER 23 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 105075-70-5 REGISTRY

ED Entered STN: 08 Nov 1986

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1-methyl-4,5-di(methyl-d3)heptyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:

CN B-Homo-7-oxaergostan-6-one-26,26,26,28,28,28-d6, 2,3,22,23-tetrahydroxy-, (2α,3α,5α,22R,23R,24S)-

FS STEREOSEARCH

MF C28 H42 D6 O6

SR CA

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT, TOXCENTER (\*File contains numerically searchable property data)

Absolute stereochemistry.

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

4 REFERENCES IN FILE CA (1907 TO DATE)

4 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 141:221056

REFERENCE 2: 139:69422

REFERENCE 3: 134:53939

REFERENCE 4: 111:233350

L7 ANSWER 24 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 93860-62-9 REGISTRY

ED Entered STN: 30 Dec 1984

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22S,23R,24S)$ - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

FS STEREOSEARCH

MF C28 H48 O6

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMINFORMRX (\*File contains numerically searchable property data)

#### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 102:19625

L7 ANSWER 25 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 93860-61-8 REGISTRY

ED Entered STN: 30 Dec 1984

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\alpha, 5\alpha, 22R, 23S)$  - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxaergostan-6-one deriv.

OTHER NAMES:

CN NSC 325611

FS STEREOSEARCH

MF C28 H48 O6

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMINFORMRX

(\*File contains numerically searchable property data)

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 102:19625

L7 ANSWER 26 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 93805-92-6 REGISTRY

ED Entered STN: 18 Dec 1984

CN B-Homo-7a-oxaergostan-7-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,225,235)$ - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5H-Benz[d]indeno[4,5-b]oxepin, B-homo-7a-oxaergostan-7-one deriv.

FS STEREOSEARCH

MF C28 H48 O6

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT

(\*File contains numerically searchable property data)

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 102:79205

L7 ANSWER 27 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 93782-67-3 REGISTRY

ED Entered STN: 18 Dec 1984

CN B-Homo-7a-oxaergostan-7-one, 2,3,22,23-tetrahydroxy-,

 $(2\alpha, 3\alpha, 5\alpha, 22R, 23R)$  - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5H-Benz[d]indeno[4,5-b]oxepin, B-homo-7a-oxaergostan-7-one deriv.

FS STEREOSEARCH

MF C28 H48 O6

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CASREACT

(\*File contains numerically searchable property data)

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 102:79205

L7 ANSWER 28 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 93518-68-4 REGISTRY

ED Entered STN: 18 Dec 1984

CN B-Homo-7-oxapregnan-6-one, 21-(2-butoxyethoxy)-2,3-dihydroxy-20-methyl-,

 $(2\alpha, 3\alpha, 5\alpha, 20S)$  - (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 6H-Benz[c]indeno[5,4-e]oxepin, B-homo-7-oxapregnan-6-one deriv.

FS STEREOSEARCH

MF C28 H48 O6

LC STN Files: CA, CAPLUS, USPATFULL

Absolute stereochemistry.

# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

2 REFERENCES IN FILE CA (1907 TO DATE)

2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 106:33373

REFERENCE 2: 102:6952

L7 ANSWER 29 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 80736-39-6 REGISTRY

ED Entered STN: 16 Nov 1984

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2S,3S,4S)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha, 3\alpha, 5\alpha, 22S, 23S, 24S)$ -

FS STEREOSEARCH

MF C28 H48 O6

CI COM

LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMCATS, CHEMINFORMRX (\*File contains numerically searchable property data)

Absolute stereochemistry.

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

8 REFERENCES IN FILE CA (1907 TO DATE)

8 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 131:130161

REFERENCE 2: 120:54778

REFERENCE 3: 116:41841

REFERENCE 4: 115:29709

REFERENCE 5: 114:116919

REFERENCE 6: 111:130875

REFERENCE 7: 97:198451

REFERENCE 8: 96:82692

L7 ANSWER 30 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN

RN 78821-43-9 REGISTRY

ED Entered STN: 16 Nov 1984

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:

```
B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,
CN
     (2\alpha, 3\alpha, 5\alpha, 22R, 23R) -
OTHER NAMES:
     24(R)-Epibrassinolide
     24-epi-Brassinolide
CN
     24-Epibrassinolide
CN
CN 24-epibrassinolide
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-(2,3-dihydroxy-1,4,5-
     trimethylhexyl)hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     [1R-[1\alpha(1S*,2R*,3R*,4R*),3a\beta,3b\alpha,6a\beta,8\beta,9\beta]
     ,10a\alpha,10b\beta,12a\alpha] ] -
CN
     B 1105
     BP 55
CN
     Epibrassinolide
CN
     Epibrassinolide R
CN
     Epin
CN
FS
     STEREOSEARCH
     126721-49-1
DR
     C28 H48 O6
MF
CI
     COM
     STN Files:
LC
                   AGRICOLA, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA, CAPLUS,
       CASREACT, CHEMCATS, CHEMINFORMRX, CSCHEM, PROMT, TOXCENTER, USPAT2,
       USPATFULL
         (*File contains numerically searchable property data)
```

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

315 REFERENCES IN FILE CA (1907 TO DATE)
5 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
315 REFERENCES IN FILE CAPLUS (1907 TO DATE).

REFERENCE 1: 143:92368

REFERENCE 2: 143:23315

REFERENCE 3: 142:350508

REFERENCE 4: 142:331198

REFERENCE 5: 142:236444

REFERENCE 6: 142:194072

REFERENCE 7: 142:192507

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REFERENCE
            8: 142:173375
REFERENCE
            9: 142:130779
REFERENCE 10: 142:110422
     ANSWER 31 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN
L7
     78821-42-8 REGISTRY
RN
     Entered STN: 16 Nov 1984
ED
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2S,3S,4R)-2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,
     (2\alpha, 3\alpha, 5\alpha, 22S, 23S) -
OTHER NAMES:
    (22S,23S)-24-Epibrassinolide
     22,23,24-Triepibrassinolide
CN
     22,23,24-Trisepibrassinolide
CN
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-(2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl)hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     [1R-[1\alpha(1S*,2S*,3S*,4R*),3a\beta,3b\alpha,6a\beta,8\beta,9\beta]
     ,10a\alpha,10b\beta,12a\alpha]]-
     B 1072
CN
CN
     Brassinosteroid
     Epibrassinolide S
CN
     Isoepibrassinolide
CN
FS
     STEREOSEARCH
     126722-25-6
DR
MF
     C28 H48 O6
CI
     COM
                   AGRICOLA, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA, CAPLUS,
LC
     STN Files:
       CASREACT, CEN, CHEMINFORMRX, CIN, PROMT, TOXCENTER, USPATZ, USPATFULL
          (*File contains numerically searchable property data)
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# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

152 REFERENCES IN FILE CA (1907 TO DATE)

17 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

152 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 143:110396

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REFERENCE
             2: 143:23106
             3: 142:424749
REFERENCE
                 142:389072
REFERENCE
REFERENCE
             5: 142:389066
             6: 142:310751
REFERENCE
                 142:236562
REFERENCE
             7:
             8: 142:236476
REFERENCE
REFERENCE
             9:
                 142:213667
REFERENCE 10: 142:33980
     ANSWER 32 OF 32 REGISTRY COPYRIGHT 2005 ACS on STN
. L7
    72962-43-7 REGISTRY
RN
     Entered STN: 16 Nov 1984
ED
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
      (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     B-Homo-7-oxaergostan-6-one, 2,3,22,23-tetrahydroxy-,
      (2\alpha, 3\alpha, 5\alpha, 22R, 23R, 24S) -
OTHER NAMES:
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-(2,3-dihydroxy-1,4,5-
     trimethylhexyl)hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
      [1R-[1\alpha(1S*,2R*,3R*,4S*),3a\beta,3b\alpha,6a\beta,8\beta,9\beta]
     ,10a\alpha,10b\beta,12a\alpha]]-
     Brassinolide
CN
     STEREOSEARCH
FS
     C28 H48 O6
MF
CI
     COM
                   AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS,
LC
     STN Files:
        BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN,
       CHEMINFORMRX, CIN, CSCHEM, EMBASE, IPA, MEDLINE, MRCK*, NAPRALERT,
        PROMT, TOXCENTER, USPAT2, USPATFULL
          (*File contains numerically searchable property data)
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\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

560 REFERENCES IN FILE CA (1907 TO DATE)
32 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
564 REFERENCES IN FILE CAPLUS (1907 TO DATE)

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REFERENCE 2: 143:148216

REFERENCE 3: 143:93950

REFERENCE 4: 143:38920

REFERENCE 5: 143:23092

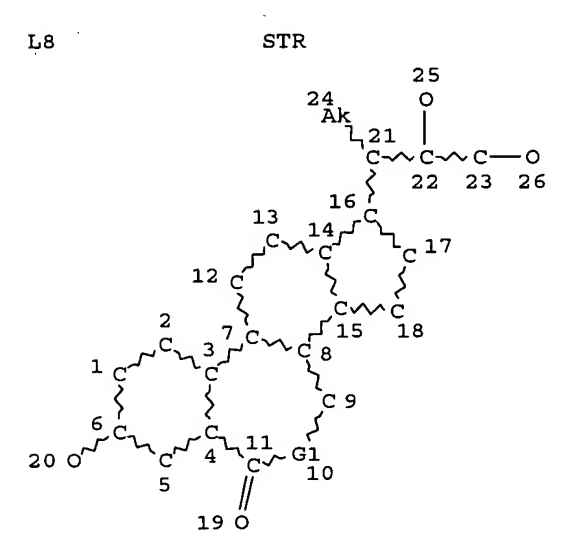
REFERENCE 6: 142:425343

REFERENCE 7: 142:352139

REFERENCE 8: 142:276838

REFERENCE 9: 142:236473

REFERENCE 10: 142:215127



REP G1=(0-1) O
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 26

STEREO ATTRIBUTES: NONE

L10 619 SEA FILE=REGISTRY SSS FUL L8

100.0% PROCESSED 9594 ITERATIONS

SEARCH TIME: 00.00.03

619 ANSWERS

=> d his full

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(FILE 'HOME' ENTERED AT 15:34:40 ON 01 SEP 2005)
     FILE 'HCAPLUS' ENTERED AT 15:34:52 ON 01 SEP 2005
              1 SEA ABB=ON PLU=ON US2004253289/PN OR US2004-711162#/AP, PRN
L1
     FILE 'REGISTRY' ENTERED AT 15:35:30 ON 01 SEP 2005
     FILE 'HCAPLUS' ENTERED AT 15:35:32 ON 01 SEP 2005
                TRA L1 1- RN:
                                      4 TERMS
L2
     FILE 'REGISTRY' ENTERED AT 15:35:32 ON 01 SEP 2005
              4 SEA ABB=ON PLU=ON L2
L3
              1 SEA ABB=ON PLU=ON L3 AND C28H48O6
L4
                D RSD
             74 SEA ABB=ON PLU=ON C5-C6-C6-OC6/ES AND C28H48O6
L5
               QUE ABB=ON PLU=ON (PMS OR MAN OR IDS OR MXS)/CI OR MIXT OR
L6
                COMPD OR COMPOUND OR UNSPECIFIED
            32 SEA ABB=ON PLU=ON L5 NOT L6
L7
                STR
L8
             38 SEA SSS SAM L8
L9
            619 SEA SSS FUL L8
L10
     FILE 'HCAPLUS' ENTERED AT 15:42:04 ON 01 SEP 2005
            924 SEA ABB=ON PLU=ON L7
L11
           1372 SEA ABB=ON PLU=ON ?EPIBRASSO? OR EPI(1A)BRASSO? OR ?BRASSINOS
L12
                TER? OR EPIN# OR NSC325611 OR NSC325(W)611 OR NSC(W)(325611 OR
                325 (W) 611) OR B1105 OR B(W) 1105 OR BP55 OR BP (W) 55
                QUE ABB=ON PLU=ON DRUG DELIVERY SYSTEMS+OLD, NT/CT
L13
                E HIV/CT
                E E3+ALL
                E E2+ALL
          47667 SEA ABB=ON PLU=ON HUMAN IMMUNODEFICIENCY VIRUS+OLD, NT/CT
L14
                E ANTI-HIV/CT
                E E4+ALL
                E E2+ALL
          22540 SEA ABB=ON PLU=ON ANTI-AIDS AGENTS+RTCS/CT
L15
                E AIDS/CT
                E E4+ALL
          17135 SEA ABB=ON PLU=ON "AIDS (DISEASE)"+OLD, NT/CT
L16
                E BRASSIN/CT
                E E4+ALL
            564 SEA ABB=ON PLU=ON BRASSINOLIDE/CT
L17
                E BRASSIN/CT
                E E5+ALL
                E HORMONES, PLANT/CT
                E E3+ALL
           1148 SEA ABB=ON PLU=ON "HORMONES, PLANT"+OLD, NT/CT (L) ?BRASSINO?
L18
L19
           1217 SEA ABB=ON PLU=ON L10
              3 SEA ABB=ON PLU=ON (L11 OR L12 OR L17 OR L18 OR L19) AND L13
L20
                E KHRIPACH V/AU
            241 SEA ABB=ON PLU=ON ("KHRIPACH V"/AU OR "KHRIPACH V A"/AU OR
L21
                "KHRIPACH V N"/AU OR "KHRIPACH V V"/AU OR "KHRIPACH VLADIMIR"/A
                U OR "KHRIPACH VLADIMIR A"/AU OR "KHRIPACH VLADIMIR V"/AU)
                E ALTSIVANOVICH/AU
              2 SEA ABB=ON PLU=ON "ALTSIVANOVICH KONSTANTIN"/AU
L22
                E ZABINSKII/AU
                                    "ZABINSKII VLADIMIR"/AU
L23
              1 SEA ABB=ON PLU=ON
                E SAMUESVICH/AU
                E SAMUSEVICH/AU
              2 SEA ABB=ON PLU=ON "SAMUSEVICH MIKHAIL"/AU
L24
                                    (DREBSK OR MIKONIK)/CS, PA
              2 SEA ABB=ON PLU=ON
L25
                E MIKONIK/CS, PA
              2 SEA ABB=ON PLU=ON L20 AND (L21 OR L22 OR L23 OR L24 OR L25)
L26
              1 SEA ABB=ON PLU=ON L20 NOT L26
L27
                                  (L11 OR L12 OR L17 OR L18 OR L19) (L) (THU
L28
             12 SEA ABB=ON PLU=ON
```

```
OR PAC OR DMA)/RL
               2 SEA ABB=ON PLU=ON L28 AND (L21 OR L22 OR L23 OR L24 OR L25)
 L29
              10 SEA ABB=ON PLU=ON L28 NOT L29
 L30
                 E HIV/CT
                 E E3+ALL
                 E HIV PROTEASE/CT
                 E E3+ALL
            4144 SEA ABB=ON PLU=ON HIV PROTEASE+NT/CT
 L31
               1 SEA ABB=ON PLU=ON (L11 OR L12 OR L17 OR L18 OR L19) AND (L14
 L32
                 OR L15 OR L16 OR L31)
               2 SEA ABB=ON PLU=ON (L26 OR L29 OR L32)

∠ L33

              13 SEA ABB=ON PLU=ON (L27 OR L28)
 L34
      FILE 'MEDLINE' ENTERED AT 16:07:30 ON 01 SEP 2005
             431 SEA ABB=ON PLU=ON (L11 OR L12 OR L19)
 L35
                 E BRASSINOLIDE/CT
                 E EPIBRASSINOLIDE/CT
                 E BRASSINOSTER/CT
                 E HIV/CT
                 E E3+ALL
           49406 SEA ABB=ON PLU=ON HIV+NT/CT
 L36
                 E E45
                 E E3+ALL
           17501 SEA ABB=ON PLU=ON REVERSE TRANSCRIPTASE INHIBITORS+NT/CT
 L37
                 E AIDS/CT
                 E E3+ALL
                 E E2
                 E E3+ALL
           63783 SEA ABB=ON PLU=ON ACQUIRED IMMUNODEFICIENCY SYNDROME/CT
 L38
                 E ANTI-HIV/CT
                 E E4+ALL
           26890 SEA ABB=ON PLU=ON ANTI-HIV AGENTS+NT/CT
 L39
                 E ANTI-AIDS/CT
                 E E4+ALL
                 E HIV PROTEASE/CT
                 E E3+ALL
            1774 SEA ABB=ON PLU=ON HIV PROTEASE/CT
 L40
                 E HIV PROTEASE INHIBITORS/CT
                 E E3+ALL
          6296 SEA ABB=ON PLU=ON HIV PROTEASE INHIBITORS+NT/CT
             200 SEA ABB=ON PLU=ON L35 AND (TU OR AD OR PD OR PK)/CT
 L42
               O SEA ABB=ON PLU=ON (L35 OR L42) AND (L36 OR L37 OR L38 OR L39
 L43
                 OR L40 OR L41)
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                 E HIV/CT
                 E E3+ALL
                 E E2+ALL
           58042 SEA ABB=ON PLU=ON HUMAN IMMUNODEFICIENCY VIRUS+NT/CT
 L44
                 E ANTI-HIV/CT
                 E E4+ALL
                 E E2+ALL
            1665 SEA ABB=ON PLU=ON ANTI HUMAN IMMUNODEFICIENCY VIRUS AGENT/CT
 L45
                 E HIV PROTEASE/CT
                 E E3+ALL
                 E E2
                 E E3+ALL
                 QUE ABB=ON PLU=ON PROTEINASE+NT/CT
 L46
                 E HIV PROTEASE INHIBITORS/CT
                 E E3+ALL
                 E E2+ALL
                 QUE ABB=ON PLU=ON PROTEINASE INHIBITOR+NT/CT
 L47
                 E AIDS/CT
                 E E3+ALL
                 E E2+ALL
           58844 SEA ABB=ON PLU=ON ACQUIRED IMMUNE DEFICIENCY SYNDROME+NT/CT
 L48
```

```
E ANTI-AIDS
               E ANTI-AIDS/CT
               E ANTI ACQUIR/CT/CT
                E ANTI ACQUIR/CT
                E ANTI-ACQUIR/CT
                                   (L11 OR L12 OR L19)
            239 SEA ABB=ON PLU=ON
L49
             31 SEA ABB=ON PLU=ON L49 AND (CB OR AD OR DT OR PD)/CT
L50
                E BRASSINLIDE/CT
                E E4+ALL
             68 SEA ABB=ON PLU=ON BRASSINOLIDE/CT
L51
                E BRASSINOSTER/CT
                E E4+ALL
            125 SEA ABB=ON PLU=ON BRASSINOSTEROID/CT
L52
             16 SEA ABB=ON PLU=ON (L51 OR L52) (L) (CB OR AD OR DT OR PD)/CT
L53
              8 SEA ABB=ON PLU=ON (L49 OR L50 OR L53) AND (L44 OR L45 OR L46
L54
                OR L47 OR L48)
             14 SEA ABB=ON PLU=ON (1999273723/AN OR 2000057269/AN OR
L55
                2001398860/AN OR 2002134469/AN OR 2002423235/AN OR 2003097079/A
               N OR 2003369601/AN OR 2004032208/AN OR 2004280593/AN OR
                2004515206/AN OR 92112219/AN OR 93307067/AN OR 94145136/AN OR
                97183315/AN) AND L50
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#### => b hcap

FILE 'HCAPLUS' ENTERED AT 16:32:46 ON 01 SEP 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 1 Sep 2005 VOL 143 ISS 10 FILE LAST UPDATED: 31 Aug 2005 (20050831/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

## => d all fhitstr 133 tot

- L33 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2005 ACS on STN
- AN 2004:1080523 HCAPLUS
- DN 142:16788
- ED Entered STN: 17 Dec 2004
- TI Natural plant compound with anti-hiv activity
- IN Khripach, Vladimir; Altsivanovich, Konstantin; Zabinskii, Vladimir; Samusevich, Mikhail
- PA Mikonik Technologies, Ltd., Belarus; Drebsk Comptech, Inc.
- SO U.S. Pat. Appl. Publ., 5 pp. CODEN: USXXCO
- DT Patent
- LA English
- IC ICM A61K031-415
  - ICS A01N043-52; A61K047-00; A61K035-78; A61K009-20; A61K009-48; A61K009-14

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INCL 424422000; 424464000; 424465000; 424439000; 424451000; 424489000;
     424725000
    1-5 (Pharmacology)
CC
     Section cross-reference(s): 11, 17
FAN. CNT 1
                                                                DATE
                                           APPLICATION NO.
     PATENT NO.
                               DATE
                        KIND
                         - - - -
                               20041216 US 2004-711162 20040828
PI US 2004253289 A1
PRAI US 2004-711162
                                20040828
CLASS
 PATENT NO.
                CLASS PATENT FAMILY CLASSIFICATION CODES
 US 2004253289 ICM
                       A61K031-415
                       A01N043-52; A61K047-00; A61K035-78; A61K009-20;
                 ICS
                       A61K009-48; A61K009-14
                       424422000; 424464000; 424465000; 424439000; 424451000;
                 INCL
                       424489000; 424725000
                       424/422.000
 US 2004253289
                 NCL
                ECLA A23L001/30; A61K031/415; A61K031/415+M; A61K045/06
    The invention comprises a method for treatment of HIV-infection and
AB
     related conditions, particularly AIDS, using plant hormone
     24-epibrassinolide, anti-HIV efficacy of which is disclosed.
    epibrassinolide natural plant hormone HIV antiHIV
ST
     Hormones, plant
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (brassinosteroids; natural plant compound, 24-
        epibrassinolide with anti-hiv activity)
     Drug delivery systems
IT
        (capsules; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
IT
     Drug delivery systems
        (coating; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     Contraceptives
IT
        (condoms; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     Drug delivery systems
        (emulsions, aqueous; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     AIDS (disease)
IT
       Anti-AIDS agents
     Combination chemotherapy
       Drug delivery systems
     Food
     Human
       Human immunodeficiency virus
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
    Natural products, pharmaceutical
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
     study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
IT
     Drug delivery systems
        (ointments, creams; natural plant compound, 24-epibrassinolide with
        anti-hiv activity)
     Drug delivery systems
IT
        (powders; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     Drug delivery systems
IT
        (solns.; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
IT
     Diet
        (supplements; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
```

```
Drug delivery systems
IT
       (suppositories, vaginal; natural plant compound, 24-epibrassinolide with
       anti-hiv activity)
    Drug delivery systems
IT
        (suspensions; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Drug delivery systems
IT
        (tablets; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    Vagina
IT
    . (tract, protection by HIV-inhibiting 24-epibrassinolid-containing composition;
       natural plant compound, 24-epibrassinolide with anti-hiv activity)
    9068-38-6
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HIV, inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    144114-21-6, HIV protease
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
       (inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    78821-43-9, 24-Epibrassinolide
IT
    RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
    52350-85-3, HIV integrase
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (of HIV, inhibitor; natural plant compound, 24-epibrassinolide with
       anti-hiv activity)
    144114-21-6, HIV protease
IT
    RL: BSU (Biological study, unclassified); BIOL (Biological study);
    PAC (Pharmacological activity); THU (Therapeutic use)
        (inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
       activity)
    144114-21-6 HCAPLUS
RN
    Retropepsin (9CI) (CA INDEX NAME)
CN
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
L33 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2005 ACS on STN
    2004:964837 HCAPLUS
AN
    141:374732
DN
    Entered STN: 12 Nov 2004
ED
    24-Epibrassinolide for decreasing cholesterol level in blood
TI
    Khripach, Vladimir; Altsivanovich, Konstantin;
IN
    Zhabinskii, Vladimir; Samusevich, Mikhail
    Mikonik Technologies, Ltd, Belarus; Drebsk Comptech,
₽A
    Inc.
SO . U.S. Pat. Appl. Publ., 6 pp.
    CODEN: USXXCO
    Patent
DT
LA English
IC ICM A61K031-365
INCL 514450000
    1-10 (Pharmacology)
    Section cross-reference(s): 11, 18, 63
FAN.CNT 1
                              DATE
                                          APPLICATION NO.
                                                                DATE
                        KIND
    PATENT NO.
                       ----
    US 2004225010 A1
                                          US 2004-710613
                                                                20040723
                              20041111
                              20040723
PRAI US 2004-710613
CLASS
 PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
 ______
 US 2004225010 ICM
                       A61K031-365
                       514450000
                INCL
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```
514/450.000
 US 2004225010
                 NCL
                        A23L001/30B2; A61K031/365
                 ECLA
     The invention discloses a method for improving blood cholesterol and its
AB
     conjugates levels in a mammal, which is based on the administration of
     steroidal plant hormone 24-epibrassinolide.
     epibrassinolide blood cholesterol plant hormone
ST
     Glycerides, biological studies
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (blood; method for decreasing cholesterol level in blood)
     Drug delivery systems
IT
        (capsules; method for decreasing cholesterol level in blood)
     Diet
IT
        (cholesterol-enriched; method for decreasing cholesterol level in
        blood)
     Drug delivery systems
IT
        (emulsions, aqueous; method for decreasing cholesterol level in blood)
    Lipoproteins
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (high-d.; method for decreasing cholesterol level in blood)
     Lipoproteins
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (low-d.; method for decreasing cholesterol level in blood)
IT
     Drug delivery systems
     Hypercholesterolemia
     Hypolipemic agents
     Nutrition, animal
        (method for decreasing cholesterol level in blood)
     Natural products, pharmaceutical
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
     study); OCCU (Occurrence); USES (Uses)
        (method for decreasing cholesterol level in blood)
     Drug delivery systems
IT
        (powders; method for decreasing cholesterol level in blood)
IT
     Drug delivery systems
        (solns.; method for decreasing cholesterol level in blood)
IT
     Diet
        (supplements; method for decreasing cholesterol level in blood)
IT
     Drug delivery systems
        (suspensions; method for decreasing cholesterol level in blood)
     Drug delivery systems
IT
        (tablets; method for decreasing cholesterol level in blood)
     57-88-5, Cholest-5-en-3-ol (3\beta)-, biological studies
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (blood; method for decreasing cholesterol level in blood)
     1406-18-4, Vitamin E 11103-57-4, Vitamin A
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (method for decreasing cholesterol level in blood)
     78821-43-9, 24-Epibrassinolide
{f IT}
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (method for decreasing cholesterol level in blood)
     78821-43-9, 24-Epibrassinolide
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
    · (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (method for decreasing cholesterol level in blood)
     78821-43-9 HCAPLUS
RN
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
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#### => d all hitstr 134 tot

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L34 ANSWER 1 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
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AN 2005:876419 HCAPLUS

Entered STN: 25 Aug 2005 ED

Antitumor application of synthetic precursors of brassinosteroids, their TI spirostane analogs, and cyclodextrin inclusion compounds

Azevedo, Mariangela De Burgos Martins; Fabrin Neto, Joao Batista; Zullo, IN Marco Antonio Teixeira; Anazetti, Maristella Conte; Quiros, Nora Marcela Haun; Melo, Patricia da Silva

Universidade Estadual de Campinas-UNICAMP, Brazil PA

Braz. Pedido PI, 16 pp. SO

CODEN: BPXXDX

Patent DT

LА Portuguese

IC ICM A61K031-724

ICS A61K035-78; A61P035-02

1-6 (Pharmacology)

Section cross-reference(s): 63

FAN.CNT 1

	PATENT NO.		KIND	DATE	APPLICATION NO.	DATE
PI	BR 20030001	33	 A	20041026	BR 2003-183	20030128
	BR 2003-183			20030128		
CLASS	ENT NO.	CLASS	PATENT	FAMILY CLASS	IFICATION CODES	
BR 2	2003000183	ICM ICS	A61K031 A61K035	-724 -78; A61P035	-02	

Synthetic precursors of 28-homobrassinosteroids, their spirostane analogs, ABand their cyclodextrin inclusion compds. may be use as antitumor agents by means of their mitochondrial dehydrogenase (MTT) reduction to induce apoptosis in leukemic HL60 cells.

brassinosteroid precursor spirostane analog antitumor  $\mathtt{ST}$ 

Animal cell line IT

(HL-60; antitumor application of synthetic precursors of brassinosteroids, their spirostane analogs, and cyclodextrin inclusion compds.)

Animal tissue culture IT

Antitumor agents

Apoptosis

Cell membrane

Human

Leukemia

(antitumor application of synthetic precursors of brassinosteroids,

```
their spirostane analogs, and cyclodextrin inclusion compds.)
     Inclusion compounds
IT
     RL: PAC (Pharmacological activity); THU (Therapeutic
     use); BIOL (Biological study); USES (Uses)
        (antitumor application of synthetic precursors of
        brassinosteroids, their spirostane analogs, and cyclodextrin
        inclusion compds.)
    Hormones, plant
IT
     RL: PAC (Pharmacological activity); THU (Therapeutic
     use); BIOL (Biological study); USES (Uses)
        (brassinosteroids; antitumor application of synthetic
        precursors of brassinosteroids, their spirostane analogs, and
        cyclodextrin inclusion compds.)
     Fibroblast
IT
        (culture of; antitumor application of synthetic precursors of
        brassinosteroids, their spirostane analogs, and cyclodextrin inclusion
        compds.)
     553-24-2, Neutral red
IT
     RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
     ANST (Analytical study); BIOL (Biological study); USES (Uses)
        (antitumor application of synthetic precursors of brassinosteroids,
        their spirostane analogs, and cyclodextrin inclusion compds.)
     9013-05-2, Phosphatase
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (antitumor application of synthetic precursors of brassinosteroids,
        their spirostane analogs, and cyclodextrin inclusion compds.)
               512-04-9, Diosgenin 4965-78-0
                                                 12619-70-4D, Cyclodextrin,
     83-48-7
IT
     inclusion compds.
                         53139-42-7
                                      58274-46-7, Isostigmasterol
     82373-95-3 83509-42-6, 28-Homocastasterone
                                                  127128-79-4
                                                             861854-06-0
     130450-01-0
                   189308-95-0
                                 189309-02-2
                                               523981-69-3
     RL: PAC (Pharmacological activity); THU (Therapeutic
     use); BIOL (Biological study); USES (Uses)
        (antitumor application of synthetic precursors of
        brassinosteroids, their spirostane analogs, and cyclodextrin
        inclusion compds.)
     9035-82-9, Dehydrogenase
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (mitochondrial MTT; antitumor application of synthetic precursors of
        brassinosteroids, their spirostane analogs, and cyclodextrin inclusion
     82373-95-3 83509-42-6, 28-Homocastasterone
IT
     RL: PAC (Pharmacological activity); THU (Therapeutic
     use); BIOL (Biological study); USES (Uses)
        (antitumor application of synthetic precursors of
       brassinosteroids, their spirostane analogs, and cyclodextrin
        inclusion compds.)
     82373-95-3 HCAPLUS
RN
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-4-ethyl-2,3-
CN
     dihydroxy-1,5-dimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
```

83509-42-6 HCAPLUS RN

Stigmastan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22R)$ CN ,23R) - (9CI) (CA INDEX NAME)

### Absolute stereochemistry.

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L34 ANSWER 2 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
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2005:427700 HCAPLUS AN

143:13202 DN

Entered STN: 20 May 2005 ED

New use of brassinolide in reversing multiple medicine resistance of tumor  ${ t TI}$ cell

Xian, Lijian; Cao, Qiyuan; Li, Yongqiang IN

Tumour Prevention and Treating Centre, Zhongshan City, Peop. Rep. China PA

Faming Zhuanli Shenqing Gongkai Shuomingshu, No pp. given SO

CODEN: CNXXEV

Patent  $\mathtt{DT}$ 

LA Chinese

IC ICM A61K031-58 ICS A61P043-00

CC 63-4 (Pharmaceuticals)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	CN 1491653	A	20040428	CN 2003-140318	20030828
PRAI	CN 2003-140318		20030828		
CLASS	5				

CLASS PATENT FAMILY CLASSIFICATION CODES PATENT NO.

```
CN 1491653
                 ICM
                        A61K031-58
                 ICS
                        A61P043-00
     The present invention relates to the new use of brassinolide in reversing
AB
     the resistance of tumor cell to multiple medicines. Brassinolide has
     powerful bioactivity, is safe and non-toxic. At very low concentration,
     brassinolide itself has no tumor inhibiting effect and can reverse the
     resistance of tumor cell with high resistance to multiple medicines.
     brassinolide drug resistance cancer anticancer agent
ST
     Multidrug resistance
IT
     Neoplasm
        (brassinolide in reversing multiple medicine resistance of tumor cell)
IT
     Hormones, plant
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (brassinosteroids; brassinolide in reversing
        multiple medicine resistance of tumor cell)
L34 ANSWER 3 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
     2004:1080523 HCAPLUS
AN
     142:16788
DN
     Entered STN: 17 Dec 2004
ED
     Natural plant compound with anti-hiv activity
TI
     Khripach, Vladimir; Altsivanovich, Konstantin; Zabinskii, Vladimir;
IN
     Samusevich, Mikhail
     Mikonik Technologies, Ltd., Belarus; Drebsk Comptech, Inc.
PA
     U.S. Pat. Appl. Publ., 5 pp.
SO
     CODEN: USXXCO
     Patent
\mathtt{DT}
     English
LA
IC
     ICM A61K031-415
     ICS A01N043-52; A61K047-00; A61K035-78; A61K009-20; A61K009-48;
          A61K009-14
INCL 424422000; 424464000; 424465000; 424439000; 424451000; 424489000;
     424725000
     1-5 (Pharmacology)
CC
     Section cross-reference(s): 11, 17
FAN.CNT 1
     PATENT NO.
                         KIND
                                DATE
                                            APPLICATION NO.
                                                                   DATE
                         ----
                                            US 2004-711162
PI US 2004253289
                       A1
                                20041216
PRAI US 2004-711162
                                20040828
CLASS
                        PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
                 CLASS
                        A61K031-415
 US 2004253289
                 ICM
                        A01N043-52; A61K047-00; A61K035-78; A61K009-20;
                 ICS
                        A61K009-48; A61K009-14
                        424422000; 424464000; 424465000; 424439000; 424451000;
                 INCL
                        424489000; 424725000
                        424/422.000
 US 2004253289
                 NCL
                 ECLA A23L001/30; A61K031/415; A61K031/415+M; A61K045/06
     The invention comprises a method for treatment of HIV-infection and
AB
     related conditions, particularly AIDS, using plant hormone
     24-epibrassinolide, anti-HIV efficacy of which is disclosed.
     epibrassinolide natural plant hormone HIV antiHIV
ST
IT
     Hormones, plant
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (brassinosteroids; natural plant compound, 24-
        epibrassinolide with anti-hiv activity)
     Drug delivery systems
ΙT
        (capsules; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     Drug delivery systems
{	t IT}
        (coating; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
```

```
IT
     Contraceptives
        (condoms; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
    Drug delivery systems
IT
        (emulsions, aqueous; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
    AIDS (disease)
IT
    Anti-AIDS agents
     Combination chemotherapy
     Drug delivery systems
     Food
     Human
     Human immunodeficiency virus
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
    Natural products, pharmaceutical
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
     study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
    Drug delivery systems
IT
        (ointments, creams; natural plant compound, 24-epibrassinolide with
        anti-hiv activity)
     Drug delivery systems
IT
        (powders; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
    Drug delivery systems
IT
        (solns.; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
IT
     Diet
        (supplements; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
    Drug delivery systems
IT
        (suppositories, vaginal; natural plant compound, 24-epibrassinolide with
        anti-hiv activity)
     Drug delivery systems
IT
        (suspensions; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     Drug delivery systems
IT
        (tablets; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
IT
     Vagina
        (tract, protection by HIV-inhibiting 24-epibrassinolid-containing composition;
       natural plant compound, 24-epibrassinolide with anti-hiv activity)
IT
     9068-38-6
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HIV, inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
    144114-21-6, HIV protease
{	t IT}
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitor; natural plant compound, 24-epibrassinolide with anti-hiv
        activity)
     78821-43-9, 24-Epibrassinolide
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
     52350-85-3, HIV integrase
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (of HIV, inhibitor; natural plant compound, 24-epibrassinolide with
        anti-hiv activity)
     78821-43-9, 24-Epibrassinolide
IT
     RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
     (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); OCCU (Occurrence); USES (Uses)
        (natural plant compound, 24-epibrassinolide with anti-hiv activity)
     78821-43-9 HCAPLUS
RN
```

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,6aS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L34 ANSWER 4 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN

AN 2004:1029441 HCAPLUS

DN 142:253719

ED Entered STN: 01 Dec 2004

TI In vitro and in vivo antiherpetic activity of three new synthetic brassinosteroid analogues

AU Michelini, Flavia M.; Ramirez, Javier A.; Berra, Alejandro; Galagovsky, Lydia R.; Alche, Laura E.

CS Laboratorio de Virologia, Depàrtamento de Quimica Biologica, Ciudad Universitaria, Facultad de Ciencias Exactas y Naturales-UBA, Buenos Aires, 1428, Argent.

SO Steroids (2004), 69(11-12), 713-720 CODEN: STEDAM; ISSN: 0039-128X

PB Elsevier B.V.

DT Journal

LA English

CC 1-5 (Pharmacology)

Brassinosteroids are a novel group of steroids that appear to be ABubiquitous in plants and are essential for normal plant growth and development. It has been previously reported that brassinosteroid analogs exert an antiviral activity against herpes simplex virus type 1 (HSV-1) and arenaviruses. In the present study, we report the chemical synthesis of compds.  $(225, 235) - 3\beta$ -bromo- $5\alpha$ , 22, 23-trihydroxystigmastan-6-one (2),  $(22S, 23S) - 5\alpha - \text{fluoro} - 3\beta - 22, 23 - \text{trihydroxystigmastan} - 6 - \text{one}$ (3),  $(22S, 23S) - 3\beta$ ,  $5\alpha$ , 22, 23-tetrahydroxy-stigmastan-6-one (4) as well as their antiherpetic activity both in a human conjunctive cell line (IOBA-NHC) and in the murine herpetic stromal keratitis (HSK) exptl. model. All compds. prevented HSV-1 multiplication in NHC cells in a dose dependent manner when added after infection with no cytotoxicity. Administration of compds. 2, 3, and 4 to the eyes of mice at 1, 2, and 3 days post-infection delayed and reduced the incidence of HSK, consisting mainly of inflammation, vascularization, and necrosis, compared to untreated, infected mice. However, viral titers of eye washes showed no differences among samples from treated and untreated mice. Since the decrease in the percentage of mice with ocular lesions occurred 5 days after treatment had ended, we suggest that brassinosteroids 2, 3, and 4 did not exert a direct antiviral effect in vivo, but rather may play a role in immune-mediated stromal inflammation, which would explain the improvement of the clin. signs of HSK observed

ST brassinosteroid analog prepn antiviral HSV1 conjunctiva keratitis

IT Antiviral agents

```
Human
     Human herpesvirus 1
        (antiherpetic activity of three new synthetic brassinosteroid analogs)
IT
     Eye
        (conjunctiva; antiherpetic activity of three new synthetic
        brassinosteroid analogs)
     Eye, disease
IT
     Inflammation
        (keratitis; antiherpetic activity of three new synthetic
        brassinosteroid analogs)
     188127-65-3P 528870-33-9P 528870-36-2P
IT
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation);
     THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (antiherpetic activity of new synthetic brassinosteroid
        analogs)
     83-48-7, Stigmasterol
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (antiherpetic activity of new synthetic brassinosteroid analogs)
     4092-62-0P
                                                167958-88-5P
                                                                188127-57-3P
                  125113-67-9P
                                 157556-31-5P
IT
                    295358-58-6P
                                   320341-60-4P
     295358-56-4P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (antiherpetic activity of new synthetic brassinosteroid analogs)
              THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
       22
RE
(1) Anon; Brassinosteroids: a new class of plant hormones 1999
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(3) Carr, J; Exp Biol Med 2001, V226, P353
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(9) Holbach, L; Ophthalmology 1990, V97, P722 MEDLINE
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(12) Liu, T; J Virol 1996, V70, P264 HCAPLUS
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(14) Ramirez, J; Steroids 2000, V65, P329 HCAPLUS
(15) Ramirez, J; Tetrahedron 2000, V56/34, P6171
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(19) Thummel, C; Genes Dev 2002, V16, P3113 HCAPLUS
(20) Wachsman, M; Antiviral Chem Chemother 2000, V11, P71 HCAPLUS
(21) Wachsman, M; Antiviral Chem Chemother 2002, V13, P61 HCAPLUS
(22) Zhao, Z; Science 1998, V279, P1344 HCAPLUS
     528870-33-9P 528870-36-2P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation);
     THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (antiherpetic activity of new synthetic brassinosteroid )
        analogs)
     528870-33-9 HCAPLUS
RN
     Stigmastan-6-one, 3,5,22,23-tetrahydroxy-, (3\beta,5\alpha,225,235)-
CN
     (9CI) (CA INDEX NAME)
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RN 528870-36-2 HCAPLUS

CN Stigmastan-6-one, 5-fluoro-3,22,23-trihydroxy-,  $(3\beta,5\alpha,225,235)$ - (9CI) (CA INDEX NAME)

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L34 ANSWER 5 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
     2004:964837 HCAPLUS
AN
    141:374732
DN
    Entered STN: 12 Nov 2004
ED
     24-Epibrassinolide for decreasing cholesterol level in blood
TI
     Khripach, Vladimir; Altsivanovich, Konstantin; Zhabinskii, Vladimir;
IN
    Samusevich, Mikhail
    Mikonik Technologies, Ltd, Belarus; Drebsk Comptech, Inc.
PA
    U.S. Pat. Appl. Publ., 6 pp.
SO
    CODEN: USXXCO
DT
   Patent
LA English
IC
    ICM A61K031-365
INCL 514450000
    1-10 (Pharmacology)
CC
     Section cross-reference(s): 11, 18, 63
FAN.CNT 1
                                                                DATE
     PATENT NO.
                        KIND
                               DATE
                                          APPLICATION NO.
                        ----
                              20041111 US 2004-710613
                                                                20040723
PI US 2004225010
                       A1
                               20040723
PRAI US 2004-710613
CLASS
 PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
```

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US 2004225010
                         A61K031-365
                  ICM
                  INCL
                         514450000
                         514/450.000
  US 2004225010
                  NCL
                         A23L001/30B2; A61K031/365
                  ECLA
      The invention discloses a method for improving blood cholesterol and its
 AB
      conjugates levels in a mammal, which is based on the administration of
      steroidal plant hormone 24-epibrassinolide.
      epibrassinolide blood cholesterol plant hormone
 ST
      Glycerides, biological studies
 IT
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (blood; method for decreasing cholesterol level in blood)
      Drug delivery systems
 IT
         (capsules; method for decreasing cholesterol level in blood)
 IT
      Diet
         (cholesterol-enriched; method for decreasing cholesterol level in
         blood)
      Drug delivery systems ·
 IT
         (emulsions, aqueous; method for decreasing cholesterol level in blood)
. IT
      Lipoproteins
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (high-d.; method for decreasing cholesterol level in blood)
      Lipoproteins
 IT
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (low-d.; method for decreasing cholesterol level in blood)
 IT
      Drug delivery systems
      Hypercholesterolemia
      Hypolipemic agents
      Nutrition, animal
         (method for decreasing cholesterol level in blood)
      Natural products, pharmaceutical
 IT
      RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
      (Pharmacological activity); THU (Therapeutic use); BIOL (Biological
      study); OCCU (Occurrence); USES (Uses)
         (method for decreasing cholesterol level in blood)
      Drug delivery systems
 IT
         (powders; method for decreasing cholesterol level in blood)
      Drug delivery systems
 IT
         (solns.; method for decreasing cholesterol level in blood)
         (supplements; method for decreasing cholesterol level in blood)
      Drug delivery systems
 IT
         (suspensions; method for decreasing cholesterol level in blood)
      Drug delivery systems
 IT
         (tablets; method for decreasing cholesterol level in blood)
      57-88-5, Cholest-5-en-3-ol (3β)-, biological studies
 IT
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (blood; method for decreasing cholesterol level in blood)
      1406-18-4, Vitamin E
                             11103-57-4, Vitamin A
 IT
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (method for decreasing cholesterol level in blood)
      78821-43-9, 24-Epibrassinolide
 IT
      RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
      (Pharmacological activity); THU (Therapeutic use); BIOL
      (Biological study); OCCU (Occurrence); USES (Uses)
         (method for decreasing cholesterol level in blood)
      78821-43-9, 24-Epibrassinolide
 IT
      RL: FFD (Food or feed use); NPO (Natural product occurrence); PAC
      (Pharmacological activity); THU (Therapeutic use); BIOL
      (Biological study); OCCU (Occurrence); USES (Uses)
         (method for decreasing cholesterol level in blood)
      78821-43-9 HCAPLUS
 RN
      6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-
 CN
      trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
      (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
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L34 ANSWER 6 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
    2004:586154 HCAPLUS
AN
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DN 141:150378

EDEntered STN: 22 Jul 2004 Inhibitors of measles virus TI

Barnard, Dale L. AU

Institute for Antiviral Research, Dept. ADVS, Utah State University, CS Logan, UT, USA

Antiviral Chemistry & Chemotherapy (2004), 15(3), 111-119 SO CODEN: ACCHEH; ISSN: 0956-3202

International Medical Press PB

Journal; General Review DT

English LA

1-0 (Pharmacology) CC

Section cross-reference(s): 15 A review. Measles virus (MV) infections have been almost eradicated in AB some industrialized nations. However, MV continues to cause severe disease and mortality in the world and is responsible for clusters of exogenous-borne disease in essentially disease-free countries. Because of the ebb and flow of immunization campaigns, especially in the poverty-stricken and war-torn Third World, and the ominous potential for severe disease and mortality, it is vital that research for discovery of therapeutic countermeasures should continue. To that end, a number of compds. have been evaluated for efficacy in vitro and in animal models, and several therapeutic modalities have been tested in the clinic. The only current therapies used in the clinic include ribavirin administered orally or i.v., alone or in combination with immune serum globulin; these therapies have demonstrated variable efficacy. Therefore, drug discovery efforts have been launched to supplement the existing treatments for MV infections. Antisense mols., adenosine and guanosine nucleosides, including ring-expanded "fat" nucleoside analogs, brassinosteroids, coumarins, peptide inhibitors, modulators of cholesterol synthesis and a variety of natural products have been screened for efficacy and toxicity both in vitro and in animals. However, none of these agents has gone into human clin. trials and most will not merit further development due to toxicity concerns and/or low potency. Thus, further research is needed to develop more potent and less toxic drugs that could be used for treating MV infections to supplement the existing MV vaccine campaigns.

review measles virus antiviral ST

Vaccines IT

(MV; inhibitors of measles virus)

Hormones, plant IT

RL: PAC (Pharmacological activity); BIOL (Biological study) (brassinosteroids; inhibitors of measles virus)

Antiviral agents IT

Human

Measles virus (inhibitors of measles virus)

IT Nucleoside analogs

RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); BIOL (Biological study)
(inhibitors of measles virus)

- IT 57-88-5, Cholesterol, biological studies
  RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); BIOL (Biological study)
- (synthesis modulators; inhibitors of measles virus)
  RE.CNT 86 THERE ARE 86 CITED REFERENCES AVAILABLE FOR THIS RECORD
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- L34 ANSWER 7 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
- AN 2004:413878 HCAPLUS
- DN 140:385420
- ED Entered STN: 21 May 2004
- TI Antiviral activity of natural and synthetic brassinosteroids
- AU Wachsman, Monica B.; Ramirez, Javier A.; Talarico, Laura B.; Galagovsky, Lydia R.; Coto, Celia E.
- CS Laboratorio de Virologia, Departamento de Quimica Biologica, Facultad de Ciencias Exactas y Naturales, Universidad de Buenos Aires, Buenos Aires, Argent.
- CODEN: CMCAFL; ISSN: 1568-0126
- PB Bentham Science Publishers Ltd.
- DT Journal; General Review
- LA English
- CC 1-0 (Pharmacology)
  - Section cross-reference(s): 11
- AB A review. Since the discovery of brassinolide, a C28 steroid with an unusual lactone B-ring structure, more than 60 related compds.

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-collectively known as brassinosteroids (BRs) - have been isolated from a
    wide variety of plant species. Exogenous application of BRs to plants at
    nanomolar to micromolar concns. has a broad spectrum of growth responses,
     such as stem elongation, inhibition of root growth, promotion of cell
    division and enhancement of stress resistance, brought about by changes in
    enzyme activity and gene expression. In the last years, biochem. and
    genetic anal. provided compelling evidence for an essential role of BRs in
    plant development. In this paper, we review our synthetic methods to
    obtain BRs analogs and report the scope of antiviral activity of these
    compds. against RNA and DNA viruses. Some of the compds. showed
     selectivity indexes (SI) 10- to 18- fold higher than ribavirin, a broad
     spectrum antiviral compound, when tested against Junin virus (JV)
     (Arenaviridae); a good antiviral activity against measles virus (MV)
     (Paramixoviridae), with SI values also higher than ribavirin used as reference
    drug, and a similar or lower activity against herpes simplex type 1 and 2
     (HSV-1 and HSV-2) (Herpesviridae) when compared to foscarnet or acyclovir,
    resp. Structure activity relationship studies (SAR) are analyzed, in
    order to detect which stereochem., type and position of functional groups
    are needed to develop a selective class of virus inhibitors.
    review antiviral natural pharmaceutical brassinosteroid structure activity
    Antiviral agents
    DNA viruses
    Human
    Human herpesvirus 1
    Human herpesvirus 2
    Junin virus
    Measles virus
     RNA viruses
        (antiviral activity of natural and synthetic brassinosteroids)
    Natural products, pharmaceutical
    RL: PAC (Pharmacological activity); THU (Therapeutic
    use); BIOL (Biological study); USES (Uses)
        (antiviral activity of natural and synthetic brassinosteroids
     Structure-activity relationship
        (antiviral; antiviral activity of natural and synthetic
       brassinosteroids)
     Hormones, plant
     RL: PAC (Pharmacological activity); THU (Therapeutic
    use); BIOL (Biological study); USES (Uses)
        (brassinosteroids; antiviral activity of natural and
       synthetic brassinosteroids)
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- L34 ANSWER 8 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
- AN 2003:888063 HCAPLUS
- DN 140:281298
- ED Entered STN: 13 Nov 2003
- On steroid part CDXVI 24-epibrassinolide at subnanomolar concentrations modulates growth and production characteristics of a mouse hybridoma
- AU Franck, Frantisek; Eckschlager, Tomas; Kohout, Ladislav
- CS Laboratory of Growth Regulators, Institute of Experimental Botany, Academy of Sciences of the Czech Republic, Prague, 102 27/10, Czech Rep.
- SO Collection of Czechoslovak Chemical Communications (2003), 68(11), 2190-2200
  - CODEN: CCCCAK; ISSN: 0010-0765
- PB Institute of Organic Chemistry and Biochemistry, Academy of Sciences of the Czech Republic
- DT Journal
- LA English
- CC 1-12 (Pharmacology)
  - Section cross-reference(s): 11, 32
- AB Brassinosteroids are known to stimulate plant growth and to possess antistress activities in plants. This work was aimed at exploring possible beneficial effects of 24-epibrassinolide on cultured mammalian

Harle 10 / 711162 cells. A mouse hybridoma was cultured either in standard serum-free medium, or in medium diluted to 30%, in which the cells underwent nutritional stress. Steady-state parameters of semicontinuous cultures conducted at 24-epibrassinolide concns. from 10-16 to 10-9 mol 1-1 were evaluated. Typical effects of the agent found both in standard and in diluted media were (i) increase in the value of mitochondrial membrane potential, (ii), drop of intracellular antibody level, (iii) increase in the fraction of the cells in the GO/G1 phase, and (iv) decrease in the fraction of the cells in the S phase. Alleviation of nutritional stress manifested itself in cultures conducted in diluted media. Viable cell d. was significantly higher (relative to control) at 24-epibrassinolide concns. 10-13 and 10-12 mol 1-1. The results of this exploratory study show that the plant hormone 24-epibrassinolide may induce perturbations in the cell division mechanism, in mitochondria performance, and in secreted protein synthesis in a mammalian cell line. At the lowest brassinosteroid concns., the number of steroid mols. in the culture was of the same order of magnitude as the number of viable cells in the culture. This implies involvement of a complex cascade mechanism, through which the steroid mol. induces alterations in gene expression leading finally to significant changes in cell culture parameters. steroid epibrassinolide antistress lymphocyte hybridoma plant growth regulator; mitochondria membrane potential cell cycle antibody brassinosteroid nutritional deprivation Antitumor agents Cell cycle Cell division Hybridoma Lymphocyte Mitosis Starvation, animal Translation, genetic (24-epibrassinolide modulates growth and production characteristics of mouse hybridoma) Interphase (cell cycle)

IT

IT

(G0-phase; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Interphase (cell cycle) IT

(G1-phase; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Interphase (cell cycle) IT

(G2-phase; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Interphase (cell cycle) ΙT

(S-phase; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Membrane potential IT

(biol.; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Hormones, plant IT

RL: PNU (Preparation, unclassified); PREP (Preparation) (brassinosteroids; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Mitochondria IT

> (membrane; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Membrane, biological IT

(mitochondrial; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Antibodies and Immunoglobulins IT

RL: BSU (Biological study, unclassified); BIOL (Biological study) (monoclonal; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

Stress, biological IT

(nutritional; 24-epibrassinolide modulates growth and production characteristics of mouse hybridoma)

78821-43-9P, 24-Epibrassinolide IT

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RL: DMA (Drug mechanism of action); PAC (Pharmacological
     activity); PRP (Properties); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (24-epibrassinolide modulates growth and production characteristics of
        mouse hybridoma)
     57-87-4, Ergosterol
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (24-epibrassinolide modulates growth and production characteristics of
        mouse hybridoma)
IT
     3037-46-5P
                  3152-46-3P
                               72050-68-1P
                                             72050-71-6P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (24-epibrassinolide modulates growth and production characteristics of
        mouse hybridoma)
RE.CNT
       26
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    78821-43-9P, 24-Epibrassinolide
     RL: DMA (Drug mechanism of action); PAC (Pharmacological
     activity); PRP (Properties); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (24-epibrassinolide modulates growth and production characteristics of
        mouse hybridoma)
     78821-43-9 HCAPLUS
RN
     6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4R)-2,3-dihydroxy-1,4,5-
CN
     trimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-,
     (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)
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Infection

HSV-1) 20817-72-5

IT

IT

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L34 ANSWER 9 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN.
     2003:590165 HCAPLUS
AN
     140:104433
DN
     Entered STN: 01 Aug 2003
ED
     Structure-activity relationship studies in a set of new brassinosteroid
TI
     derivatives assayed against herpes simplex virus type 1 and 2 in cell
     cultures
     Talarico, Laura B.; Ramirez, Javier A.; Galagovsky, Lydia R.; Wachsman,
ΑU
     Monica B.
     Laboratorio de Virologia. Departamento de Quimica Biologica, Universidad
CS
     de Buenos Aires, Ciudad Universitaria, Pabellon 2, Piso 4, Buenos Aires,
     1428, Argent.
     Medicinal Chemistry Research (2002), 11(8), 434-444
SO
     CODEN: MCREEB; ISSN: 1054-2523
     Birkhaeuser Boston
ΡB
     Journal /
DT
     English
LА
     1-3 (Pharmacology)
CC
     Section cross-reference(s): 2
     Thirty-seven brassinosteroid derivs. were tested for their antiviral
AB
     activity against herpes simplex virus (HSV) type 1 and twenty-seven
     against HSV type 2, via a virus yield reduction assay. Most of the assayed
     compds. show selectivity indexes (SI) higher than those obtained with the
     reference drug, stigmasterol. The compds. that possessed a better
     structure-activity relationship are 6b [(22S,23S)-3β-bromo-
     5\alpha, 22, 23-trihydroxystigmastan-6-one], 7b [(22S, 23S)-
     3\beta, 5\alpha, 22, 23-tetrahydroxystigmastan-6-one] and 12b
     [(225, 235) - 5\alpha - \text{fluor} - 3\beta, 22, 23 - \text{trihydroxy} - \text{stigmastan} - 6 - \text{one}] with
     SI values of 100, 80 and 109 for HSV-1 and 71, 40 and 27 for HSV-2, resp.
     brassinosteroid analog antiviral structure herpes simplex virus
ST
     Structure-activity relationship
IT
         (HSV inhibiting; structure-activity of brassinosteroid derivs. against
        HSV-1 and HSV-1)
     Structure-activity relationship
IT
         (antiviral; structure-activity of brassinosteroid derivs. against HSV-1
        and HSV-1)
     Antiviral agents
IT
     Human
     Human herpesvirus 1
     Human herpesvirus 2
```

(structure-activity of brassinosteroid derivs. against HSV-1 and HSV-1)

(viral; structure-activity of brassinosteroid derivs. against HSV-1 and

81481-12-1 83509-42-6 83510-06-9

85197-40-6 90524-90-6 90524-93-9 135158-75-7

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157556-31-5 174656-45-2
    147200-28-0
    188127-43-7 188127-46-0 188127-49-3
                                             188127-52-8
    188127-57-3 188127-61-9 188127-63-1 188127-64-2
    188127-65-3 220845-39-6 295358-52-0 295358-54-2
    301699-56-9 398143-21-0
                                 398143-22-1
                                               528870-32-8 528870-33-9
    528870-34-0 528870-35-1 528870-36-2
                                             528870-37-3
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    646522-48-7 646522-49-8
    RL: PAC (Pharmacological activity); PRP (Properties); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (structure-activity of brassinosteroid derivs. against HSV-1
       and HSV-1)
             THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD
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IT
     90524-93-9 135158-75-7 147200-28-0
     174656-45-2 188127-43-7 188127-46-0
     188127-61-9 188127-63-1 295358-52-0
     295358-54-2 528870-33-9 528870-36-2
     646522-46-5 646522-47-6 646522-48-7
     646522-49-8
     RL: PAC (Pharmacological activity); PRP (Properties); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (structure-activity of brassinosteroid derivs. against HSV-1
        and HSV-1)
     83509-42-6 HCAPLUS
RN
     Stigmastan-6-one, 2,3,22,23-tetrahydroxy-, (2\alpha,3\alpha,5\alpha,22R)
CN
     ,23R) - (9CI) (CA INDEX NAME)
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RN 83510-06-9 HCAPLUS CN Stigmastan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22S$ ,23S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

Absolute stereochemistry.

RN 90524-93-9 HCAPLUS

CN Stigmastan-6-one, 3,22,23-trihydroxy-,  $(3\alpha,5\alpha,22R,23R)$ - (9CI)

(CA INDEX NAME)

Absolute stereochemistry.

RN 135158-75-7 HCAPLUS

CN Stigmastan-6-one, 2,3,5,22,23-pentahydroxy-,  $(2\alpha,3\alpha,5\alpha,2)$ 2S,23S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 147200-28-0 HCAPLUS

CN 6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-4-ethyl-2,3-dihydroxy-1,5-dimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R,3aS,3bS,8S,9R,10aR,10bS,12aS)- (9CI) (CA INDEX NAME)

RN 174656-45-2 HCAPLUS CN Stigmastan-6-one, 2,3,5,22,23-pentahydroxy-,  $(2\alpha,3\alpha,5\alpha,2)$  2R,23R)- (9CI) (CA INDEX NAME)

## Absolute stereochemistry.

RN 188127-43-7 HCAPLUS CN Stigmastan-6-one, 3-(acetyloxy)-22,23-dihydroxy-,  $(3\beta,5\alpha,22R,23R)$ - (9CI) (CA INDEX NAME)

## Absolute stereochemistry.

RN 188127-46-0 HCAPLUS

CN Stigmastan-6-one, 3-(acetyloxy)-5,22,23-trihydroxy-,  $(3\beta, 5\alpha, 22R, 23R)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-61-9 HCAPLUS CN Stigmastan-6-one, 3-(acetyloxy)-22,23-dihydroxy-,  $(3\beta,5\alpha,225,235)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-63-1 HCAPLUS CN Stigmastan-6-one, 3-(acetyloxy)-5,22,23-trihydroxy-,  $(3\beta,5\alpha,225,235)$ - (9CI) (CA INDEX NAME)

RN 295358-52-0 HCAPLUS
CN Stigmastan-6-one, 3,5,22,23-tetrahydroxy-, (3β,5α,22R,23R)(9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

RN 528870-33-9 HCAPLUS

CN Stigmastan-6-one, 3,5,22,23-tetrahydroxy-,  $(3\beta,5\alpha,22S,23S)$ -

# (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 528870-36-2 HCAPLUS

CN Stigmastan-6-one, 5-fluoro-3,22,23-trihydroxy-, (3 $\beta$ ,5 $\alpha$ ,22S,23S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 646522-46-5 HCAPLUS

CN Stigmastan-6-one, 5-fluoro-2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22S,23S)$ - (9CI) (CA INDEX NAME)

RN 646522-47-6 HCAPLUS

CN Stigmastan-6-one, 3,5,22,23-tetrahydroxy-,  $(3\alpha,5\alpha,22S,23S)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 646522-48-7 HCAPLUS

CN Stigmast-4-ene-3,6-dione, 22,23-dihydroxy-, (22R,23R)- (9CI) (CA INDEX NAME).

Absolute stereochemistry.

RN 646522-49-8 HCAPLUS

CN Stigmast-4-ene-3,6-dione, 22,23-dihydroxy-, (225,235)- (9CI) (CA INDEX NAME)

ANSWER 10 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN L34 2002:829293 HCAPLUS AN 138:395464 DN Entered STN: 31 Oct 2002 ED Antiviral activity of brassinosteroids derivatives against measles virus TI in cell cultures Wachsman, Monica B.; Ramirez, Javier A.; Galagovsky, Lydia R.; Coto, Celia AU Ε. Laboratorio de Virologia, Departamento de Quimica Biologica, Universidad CS de Buenos Aires, Buenos Aires, Argent. Antiviral Chemistry & Chemotherapy (2002), 13(1), 61-66 SO CODEN: ACCHEH; ISSN: 0956-3202 International Medical Press PB Journal DTEnglish LA1-5 (Pharmacology) CCTwenty-seven brassinosteroid derivs. were tested for antiviral activity AB against measles virus (MV) via a virus-yield reduction assay. Compds. [ $(22S, 23S) - 3\beta$ -bromo- $5\alpha$ , 22, 23-trihydroxystigmastan-6-one], [ $(22R, 23R) - 2\alpha, 3\alpha, 22, 23$ -tetrahydroxy- $\beta$ -Homo-7-oxastigmastan-6-one], [(22R, 23R)-3β-fluoro-22, 23-dihydroxystigmastan-6one],  $[(225, 235) - 3\beta$ -fluoro- $5\alpha$ , 22, 23-trihydroxystigmastan-6-one] and  $[(22S, 23S) - 5\alpha - \text{fluor} - 3\beta, 22, 23 - \text{trihydroxystigmastan} - 6 - \text{one}]$ , are the derivs. with good antiviral activity against MV. These SI values are higher than those obtained with ribavirin (used as reference drug). A comparative anal. of 50% cytotoxic concentration (CC50) values, using confluent non-growing cells, gives and indication of structure-activity relationship. According to their degree of cytotoxicity the compds. were divided in three groups: low, intermediate and high cytotoxicity. By observing the chemical structures of compds. belonging to the first group we can see that less cytotoxic activities are related to the presence of a 3β-hydroxy group on C-3 (ring A) and a double bond between C-22 and C-23 (side chain). The replacement of a  $5\alpha$ -hydroxy group by a  $5\alpha$ -fluoro group enhances cytotoxicity. Halogenated brassinosteroid derivs. in C-3 position are more cytotoxic than those with an acetoxy group in the same position. For 3 compds. and ribavirin, cytotoxicity measurements were also done with replicating cells; CC50 values were low,

ST brassinosteroid antiviral measles virus

IT Antiviral agents

Measles virus

(antiviral activity of brassinosteroids derivs. against measles virus in cell cultures)

IT Hormones, plant

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (brassinosteroids; antiviral activity of

but they still competed favorably with ribavirin against MV.

brassinosteroids derivs. against measles virus in cell cultures) IT 83-48-7, Stigmasterol 36791-04-5, Ribavirin 81481-12-1 82373-95-3 83509-42-6 83510-06-9 135158-75-7 157556-31-5 174656-45-2 188127-43-7 188127-46-0 188127-49-3 188127-52-8 188127-57-3 **188127-61-9 188127-63-1** 188127-64-2 188127-65-3 295358-52-0 295358-54-2 301699-56-9 528870-34-0 398143-22-1 528870-32-8 **528870-33-9** 528870-37-3 528870-72-6 528870-35-1 **528870-36-2** RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antiviral activity of brassinosteroids derivs. against measles virus in cell cultures) THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 16 RE (1) Banks, G; Clinical applications of ribavirin 1980, P203 (2) Barnard, D; Antiviral Chemistry & Chemotherapy 2001, V12, P241 HCAPLUS (3) Cutts, F; Vaccine 1999, V17, PS47 (4) De Clerq, E; Antimicrobial Agents and Chemotherapy 1991, V35, P679 (5) Denizot, F; Journal of Immunological Methods 1986, V89, P271 MEDLINE (6) Forni, A; Clinical Infectious Disease 1994, V19, P454 MEDLINE (7) Gururangen, S; Journal of Infectious Disease 1990, V20, P219 (8) McMorris, T; Phytochemistry 1994, V36, P585 HCAPLUS (9) Ramirez, J; Steroids 2000, V65, P329 HCAPLUS (10) Ramirez, J; Tetrahedron 2000, V56, P6171 HCAPLUS (11) Ross, L; American Journal of Medicine 1984, V88, P313 (12) Shigeta, S; Antimicrobial Agents and Chemotherapy 1992, V36, P435 HCAPLUS (13) Stogner, S; Southern Medical Journal 1993, V141, P1415 (14) Teme Centurion, O; Anales de la Asociacion Quimica Argentina 1998, V86, P104 HCAPLUS (15) Wachsman, M; Antiviral Chemistry & Chemotherapy 2000, V11, P71 HCAPLUS (16) Wyde, P; Antmicrobial Agents and Chemotherapy 2000, V44, P1146 HCAPLUS 82373-95-3 83509-42-6 83510-06-9 135158-75-7 174656-45-2 188127-43-7 18812 7-46-0 188127-61-9 188127-63-1 295358-52-0 295358-54-2 528870-33-9 528870-36-2 RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antiviral activity of brassinosteroids derivs. against measles virus in cell cultures) 82373-95-3 HCAPLUS RN6H-Benz[c]indeno[5,4-e]oxepin-6-one, 1-[(1S,2R,3R,4S)-4-ethyl-2,3-CN dihydroxy-1,5-dimethylhexyl]hexadecahydro-8,9-dihydroxy-10a,12a-dimethyl-, (1R, 3aS, 3bS, 6aS, 8S, 9R, 10aR, 10bS, 12aS) - (9CI) (CA INDEX NAME)

RN 83509-42-6 HCAPLUS

CN Stigmastan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22R,23R)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 83510-06-9 HCAPLUS

CN Stigmastan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22S,23S)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 135158-75-7 HCAPLUS

CN Stigmastan-6-one, 2,3,5,22,23-pentahydroxy-,  $(2\alpha,3\alpha,5\alpha,2)$  2S,23S)- (9CI) (CA INDEX NAME)

RN 174656-45-2 HCAPLUS

CN Stigmastan-6-one, 2,3,5,22,23-pentahydroxy-,  $(2\alpha,3\alpha,5\alpha,22,23)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-43-7 HCAPLUS

CN Stigmastan-6-one, 3-(acetyloxy)-22,23-dihydroxy-,  $(3\beta, 5\alpha, 22R, 23R)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-46-0 HCAPLUS

CN Stigmastan-6-one, 3-(acetyloxy)-5,22,23-trihydroxy-,

 $(3\beta, 5\alpha, 22R, 23R)$  - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-61-9 HCAPLUS CN Stigmastan-6-one, 3-(acetyloxy)-22,23-dihydroxy-,  $(3\beta,5\alpha,22S,23S)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-63-1 HCAPLUS CN Stigmastan-6-one, 3-(acetyloxy)-5,22,23-trihydroxy-,  $(3\beta,5\alpha,225,235)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN  $_{\circ}$  295358-54-2 HCAPLUS CN Stigmastan-6-one, 5-fluoro-3,22,23-trihydroxy-, (3 $\beta$ ,5 $\alpha$ ,22R,23R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 528870-33-9 HCAPLUS CN Stigmastan-6-one, 3,5,22,23-tetrahydroxy-,  $(3\beta,5\alpha,22S,23S)$ - (9CI) (CA INDEX NAME)

528870-36-2 HCAPLUS RN

Stigmastan-6-one, 5-fluoro-3,22,23-trihydroxy-,  $(3\beta,5\alpha,225,235)$ -CN (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L34 ANSWER 11 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN

AN 2000:129185 HCAPLUS

132:273847 DN

Entered STN: 25 Feb 2000 ED

Antiviral effect of brassinosteroids against herpes virus and arenaviruses TI

Wachsman, Monica B.; Lopez, Elsa M. F.; Ramirez, Javier A.; Galagovsky, AU

Lydia R.; Coto, Celia E.

Laboratorio de Virologia, Departamento de Quimica Biologica and Facultad CS de Ciencias Exactas y Naturales, Universidad de Buenos Aires, Buenos Aires, 1428, Argent.

Antiviral Chemistry & Chemotherapy (2000), 11(1), 71-77 SO CODEN: ACCHEH; ISSN: 0956-3202

International Medical Press ΡB

Journal DT

English LΑ

1-3 (Pharmacology) CC

Section cross-reference(s): 11

A natural brassinosteroid and a series of synthetic derivs. were found to ABbe good inhibitors of herpes simplex virus type 1 (HSV-1) and arenavirus replication in cell culture. The synthetic compds. tested were analogs of the 24(S) ethylbrassinone. Compds.  $(22R, 23R, 24S)-2\alpha$ ,  $3\alpha, 5\alpha, 22, 23$ -pentahydroxy-stigmastan-6-one and

 $(22R, 23R, 24S) - 3\beta$ -bromo- $5\alpha$ , 22, 23-trihydroxy-stigmastan-6-one were cytotoxic at concns. of 20-40  $\mu$ M. (22S,23S,24S)-

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2\alpha, 3\alpha, 22, 23-tetrahydroxy-5\alpha, stigmastan-6-one,
     (22R, 23R, 24S) - 3\beta-acetoxy-22,23-dihydroxy-5\alpha-cholestan-6-one,
     (22S, 23S, 24S) - 3\beta-bromo-22, 23-dihydroxy-5\alpha-chol-estan-6-one and
     (22S, 23S, 24S) - 3\beta-bromo-5\alpha, 22, 23-trihydroxy-stigmastan-6-one
    were the most active of the series against HSV-1, with selectivity index
     (SI) values (CC50/EC50) ranging from 10.6 to 16.5. The majority of the
    compds. were potent inhibitors of arenaviruses, (22S, 23S, 24S) - 3\beta-
    bromo-5\alpha,22,23-trihydroxy-stig-mastan-6-one being the most active,
    with SI values of 307.8 and 692.5 for Tacaribe and Junin viruses, resp.
    The antiviral activity of brassinosteroid derivs. was not because of
    direct inactivation; time-of-addition expts. suggested that a late step in
    HSV-1 multiplication was affected, whereas arenaviruses remained
    susceptible to the compds. throughout the replicative cycle.
    natural brassinosteroid antiviral SAR HSV1; arenavirus Junin virus
    inhibiting brassinosteroid structure
    Antiviral agents
    Arenavirus
     Human herpesvirus 1
    Structure-activity relationship
        (antiviral effect of brassinosteroids against HSV and arenaviruses)
    Natural products, pharmaceutical
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); PRP (Properties); THU (Therapeutic use);
     BIOL (Biological study); USES (Uses)
        (antiviral effect of brassinosteroids against HSV and
        arenaviruses)
    Hormones, plant
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); PRP (Properties); THU (Therapeutic use);
     BIOL (Biological study); USES (Uses)
        (brassinosteroids; antiviral effect of
       brassinosteroids against HSV and arenaviruses)
     83509-42-6 83510-06-9 135158-75-7
     174656-45-2 188127-43-7 188127-46-0
     188127-49-3 188127-52-8 188127-61-9 188127-63-1
     188127-64-2 188127-65-3
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); PRP (Properties); THU (Therapeutic use);
     BIOL (Biological study); USES (Uses)
        (antiviral effect of brassinosteroids against HSV and
        arenaviruses)
              THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD
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(20) McMorris, T; Phytochemistry 1994, V36, P585 HCAPLUS
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Absolute stereochemistry.

RN 83510-06-9 HCAPLUS CN Stigmastan-6-one, 2,3,22,23-tetrahydroxy-,  $(2\alpha,3\alpha,5\alpha,22S,23S)$  - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 135158-75-7 HCAPLUS CN Stigmastan-6-one, 2,3,5,22,23-pentahydroxy-,  $(2\alpha,3\alpha,5\alpha,2)$  2S,23S)- (9CI) (CA INDEX NAME)

RN 174656-45-2 HCAPLUS

CN Stigmastan-6-one, 2,3,5,22,23-pentahydroxy-,  $(2\alpha,3\alpha,5\alpha,2)$  2R,23R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-43-7 HCAPLUS

CN Stigmastan-6-one, 3-(acetyloxy)-22,23-dihydroxy-,  $(3\beta,5\alpha,22R,23R)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-46-0 HCAPLUS

CN Stigmastan-6-one, 3-(acetyloxy)-5,22,23-trihydroxy-,

 $(3\beta, 5\alpha, 22R, 23R)$  - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-61-9 HCAPLUS
CN Stigmastan-6-one, 3-(acetyloxy)-22,23-dihydroxy-,

 $(3\beta, 5\alpha, 22S, 23S) - (9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

RN 188127-63-1 HCAPLUS CN Stigmastan-6-one, 3-(acetyloxy)-5,22,23-trihydroxy-,  $(3\beta,5\alpha,225,235)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

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L34 ANSWER 12 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
     1999:27929 HCAPLUS
AN
     130:91278
DN
     Entered STN: 14 Jan 1999
ED
     Steroid receptor kinase BIN1 involved in brassinosteroid signal
TI
     transduction from Arabidopsis thaliana
     Chory, Joanne; Li, Jianming
IN
     The Salk Institute for Biological Studies, USA
PA
SO
     PCT Int. Appl., 72 pp.
     CODEN: PIXXD2
DT
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     ICM C12N005-00
     ICS C12N015-00; C07H021-02; C12Q001-00; C07K001-00; C07K016-00
     3-3 (Biochemical Genetics)
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     Section cross-reference(s): 1, 6, 11
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             UA, UG, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
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             CM, GA, GN, ML, MR, NE, SN, TD, TG
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CLASS
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 PATENT NO.
                 CLASS
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 WO 9859039
                 ICM
                        C12N015-00; C07H021-02; C12Q001-00; C07K001-00;
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                        C07K014/415; C12N015/82C8; G01N033/74B
 WO 9859039
                 ECLA
                        800/290.000; 435/007.100; 435/007.800; 435/069.100;
 US 6245969
                 NCL
                        435/194.000; 435/320.100; 435/419.000; 435/421.000;
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                        800/279.000; 800/286.000; 800/301.000
                        C07K014/415; C12N015/82C8; G01N033/74B
                 ECLA
     A novel plant steroid receptor, Bin1, is provided, as well as
AB
     polynucleotides encoding Bin1. Bin1 polypeptide is useful in promoting
     increased plant yield and/or increased plant biomass. Arabidopsis dwarf
     mutants were identified that were unable to respond to exogenously added
     brassinosteroid, a phenotype that might be expected for brassinosteroid
     signaling mutants. All mutations defined alleles of a single previously
     described gene, BRI1. BRI1 was cloned and its expression pattern examined
     It encodes a ubiquitously expressed putative receptor kinase. The
     extracellular domain contains 25 tandem leucine-rich repeats that resemble
     repeats found in animal hormone receptors, plant disease resistance genes,
     and genes involved in unknown signaling pathways controlling plant
     development. Thus, genetically modified plants characterized as having
     increased yield and methods for producing such plants are provided, as are
```

```
transgenic animals in which oocyte maturation is stimulated.
ST
    Arabidopsis receptor kinase BIN1 cDNA sequence; brassinosteroid signal
     transduction Arabidopsis receptor kinase
     Chromosome
IT
        (Arabidopsis thaliana 4, gene mapping on; steroid receptor kinase BIN1
        involved in brassinosteroid signal transduction from Arabidopsis
        thaliana)
    Gene, plant
IT
     RL: AGR (Agricultural use); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); USES (Uses)
        (BRI1; steroid receptor kinase BIN1 involved in brassinosteroid
        signal transduction from Arabidopsis thaliana)
     Steroid receptors
IT
    RL: AGR (Agricultural use); BAC (Biological activity or effector, except
     adverse); BSU (Biological study, unclassified); PRP (Properties); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (Bin1; steroid receptor kinase BIN1 involved in brassinosteroid
        signal transduction from Arabidopsis thaliana)
     Promoter (genetic element)
IT
     RL: AGR (Agricultural use); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (FMV35S or CaMV35S or pathogen infection-induced; steroid receptor
        kinase BIN1 involved in brassinosteroid signal transduction
        from Arabidopsis thaliana)
    Hormones, plant
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (brassinosteroids; steroid receptor kinase BIN1 involved in
        brassinosteroid signal transduction from Arabidopsis thaliana)
     cDNA sequences
IT
        (for leucine-rich repeat steroid receptor kinase Bin1 from Arabidopsis
        thaliana)
     Genetic mapping
IT
        (gene mapping on Arabidopsis chromosome 4; steroid receptor kinase BIN1
        involved in brassinosteroid signal transduction from Arabidopsis
        thaliana)
    Disease resistance, plant
IT
     Oogenesis
        (genetic engineering for; steroid receptor kinase BIN1 involved in
        brassinosteroid signal transduction from Arabidopsis thaliana)
     Animal cell
IT
        (mammalian, transgenic; steroid receptor kinase BIN1 involved in
        brassinosteroid signal transduction from Arabidopsis thaliana)
IT
     Protein sequences
        (of leucine-rich repeat steroid receptor kinase Bin1 from Arabidopsis
        thaliana)
    Arabidopsis thaliana
IT
     Genetic engineering
     Signal transduction, biological
        (steroid receptor kinase BIN1 involved in brassinosteroid signal
        transduction from Arabidopsis thaliana)
    Antisense DNA
IT
     RL: AGR (Agricultural use); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (steroid receptor kinase BIN1 involved in brassinosteroid
        signal transduction from Arabidopsis thaliana)
     Antibodies
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (steroid receptor kinase BIN1 involved in brassinosteroid signal
        transduction from Arabidopsis thaliana)
     Plant cell
IT
     Seed
        (transgenic; steroid receptor kinase BIN1 involved in brassinosteroid
        signal transduction from Arabidopsis thaliana)
                                 219315-26-1
     197181-05-8
                   219306-79-3
IT
     RL: AGR (Agricultural use); BAC (Biological activity or effector, except
     adverse); BSU (Biological study, unclassified); PRP (Properties); THU
```

```
(Therapeutic use); BIOL (Biological study); USES (Uses)
        (amino acid sequence; steroid receptor kinase BIN1 involved in
        brassinosteroid signal transduction from Arabidopsis thaliana)
     196526-86-0, GenBank AF017056
IT
     RL: AGR (Agricultural use); PRP (Properties); THU (Therapeutic
     use); BIOL (Biological study); USES (Uses)
        (nucleotide sequence; steroid receptor kinase BIN1 involved in
        brassinosteroid signal transduction from Arabidopsis thaliana)
              THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 1
RE
(1) Li; Cell 1997, V90, P929 HCAPLUS
L34 ANSWER 13 OF 13 HCAPLUS COPYRIGHT 2005 ACS on STN
AN
     1936:63968 HCAPLUS
DN
     30:63968
OREF 30:8527b-d
     Entered STN: 16 Dec 2001
ED
     Some new compounds of hexamethylenetetramine
TI
     Bouchereau, P.
AU
     Journal de Pharmacie et de Chimie (1936), 23, 549-56
SO
     CODEN: JPHCA9; ISSN: 0368-3591
DT
     Journal
     Unavailable
LA
     17 (Pharmaceuticals, Cosmetics, and Perfumes)
CC
     Neutral or feebly alkaline derivs. of C6H12N4 (X), e.g., diphenate (cf. B. in
AB
     Douris and Beytout, C. A. 17, 1621) are no longer caustic, toxicity is
     greatly lessened and the bactericidal or other action of, e.g., the
     phenol constituent is much increased; i. e., the action of the complexes
     is sp. (CaCl2) X.2H2O, crystalline, is formed by precipitation of concentrated
solution of X with
     hot CaCl2 solution; at 75-80°, HCHO is given off without m.; it is
     soluble in H2O, little soluble in alc., insol. in Et2O. Assay methods are
     given; an accurate method of determining X is based on precipitating the nearly insol.
     compound (HgCl2)2X.H2O (Del.acte.epine). The CaCl2 compound is an
     active diuretic, hemostatic and recalcifiant in pulmonary tuberculosis.
     (MqCl2)2X.H2O is soluble in 5.5 parts H2O at 15°; it is antiseptic, a
     sedative in liver troubles and a wound antiseptic. (MgS203)2X.H2O,
     crystalline, is soluble in H2O. In the assay for X by HgCl2, a brown color of the
     precipitate causes a slight error.
     Pharmaceutical preparations
{	t IT}
        (hexamethylenetetramine compds.)
     Calcium chloride, compound with hexamethylenetetramine
IT
     Magnesium thiosulfate, compound with hexamethylenetetramine
     100-97-0, Hexamethylenetetramine
IT
        (compds. of)
     4015-89-8, Mercury chloride, HgCl2, compound with hexamethylenetetramine
IT
     859193-53-6, Magnesium chloride, compound with hexamethylenetetramine
        (preparation of)
=> b embase
FILE 'EMBASE' ENTERED AT 16:33:09 ON 01 SEP 2005
COPYRIGHT (C) 2005 Elsevier Inc. All rights reserved.
 FILE COVERS 1974 TO 25 Aug 2005 (20050825/ED)
 EMBASE has been reloaded. Enter HELP RLOAD for details.
 This file contains CAS Registry Numbers for easy and accurate
 substance identification.
=> d all 155 tot
L55 ANSWER 1 OF 14 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.
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on STN

2004515206 EMBASE

AN

```
In vitro and in vivo antiherpetic activity of three new synthetic
TI
     brassinosteroid analogues.
     Michelini F.M.; Ramirez J.A.; Berra A.; Galagovsky L.R.; Alche L.E.
AU
     lalche@qb.fcen.uba.ar
CS
     Steroids, (2004) Vol. 69, No. 11-12, pp. 713-720.
SO
     Refs: 22
     ISSN: 0039-128X CODEN: STEDAM
PUI
     S 0039-128X(04)00141-2
CY
     United States
     Journal; Article
DT
             Microbiology
FS
     004
             Pharmacology
     030
             Drug Literature Index
     037
LA
     English
     English
SL
     Entered STN: 20041230
ED
     Last Updated on STN: 20041230
     Brassinosteroids are a novel group of steroids that appear to be
AB
     ubiquitous in plants and are essential for normal plant growth and
     development. It has been previously reported that brassinosteroid
     analogues exert an antiviral activity against herpes simplex virus type 1
     (HSV-1) and arenaviruses. In the present study, we report the chemical
     synthesis of compounds (22S, 23S) - 3\beta-bromo-5\alpha, 22, 23-
     trihydroxystigmastan-6-one (2), (22S, 23S)-5\alpha-fluoro-3\beta-22, 23-6
     trihydroxystigmastan-6-one (3), (22S, 23S)-3\beta, 5\alpha, 22, 23-
     tetrahydroxy-stigmastan-6-one (4) as well as their antiherpetic activity
     both in a human conjunctive cell line (IOBA-NHC) and in the murine
     herpetic stromal keratitis (HSK) experimental model. All compounds
     prevented HSV-1 multiplication in NHC cells in a dose dependent manner
     when added after infection with no cytotoxicity. Administration of
     compounds 2, 3, and 4 to the eyes of mice at 1, 2, and 3 days
     post-infection delayed and reduced the incidence of HSK, consisting mainly
     of inflammation, vascularization, and necrosis, compared to untreated,
     infected mice. However, viral titers of eye washes showed no differences
     among samples from treated and untreated mice. Since the decrease in the
     percentage of mice with ocular lesions occurred 5 days after treatment had
     ended, we suggest that brassinosteroids 2, 3, and 4 did not
     exert a direct antiviral effect in vivo, but rather may play a role in
     immune-mediated stromal inflammation, which would explain the improvement
     of the clinical signs of HSK observed. .COPYRGT. 2004 Elsevier Inc. All
     rights reserved.
     Medical Descriptors:
CT
     *drug synthesis
     in vitro study
     in vivo study
     stomatitis
     antiviral activity
     cell line
     cytotoxicity
     Herpes simplex virus 1
     virus infection: ET, etiology
     inflammation
     vascularization
     necrosis
     comparative study
     eye injury
     sample
     morbidity
     dose response
     cell division
     immune mediated injury
     human
     nonhuman
     male
     mouse
     human cell
```

```
animal cell
     article
     Drug Descriptors:
       *brassinosteroid: PD, pharmacology
       3beta bromo 5alpha 22,23 trihydroxystigmastan 6 one: PD,
     pharmacology
       5alpha fluoro 3beta 22,23 trihydroxystigmastan 6 one: PD,
     pharmacology
       3beta, 5alpha, 22, 23 tetrahydroxystigmastan 6 one: PD, pharmacology
       antivirus agent: PD, pharmacology
       steroid: PD, pharmacology
     unclassified drug
L55 ANSWER 2 OF 14 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.
     on STN
     2004280593 EMBASE
AN
     Antiviral activity of natural and synthetic brassinosteroids.
TI
     Wachsman M.B.; Ramirez J.A.; Talarico L.B.; Galagovsky L.R.; Coto C.E.
AU
     M.B. Wachsman, Laboratorio de Virologia, Departamento de Quimica
CS
     Biologica, Universidad de Buenos Aires, Pabellon 2, Piso 4, 1428 Buenos
     Aires, Argentina. wachsman@qb.fcen.uba.ar
     Current Medicinal Chemistry: Anti-Infective Agents, (2004) Vol. 3, No. 2,
SO
     pp. 163-179.
     Refs: 64
     ISSN: 1568-0126 CODEN: CMCAFL
     Netherlands
CY
     Journal; General Review
DT
             Microbiology
FS
     004
     030
             Pharmacology
             Drug Literature Index
     037
     English
LΑ
     English
\mathtt{SL}
     Entered STN: 20040722
ED
     Last Updated on STN: 20040722
     Since the discovery of brassinolide, a C(28) steroid with an unusual
AB
     lactone B-ring structure, more than 60 related compounds -collectively
     known as brassinosteroids (BRs) - have been isolated from a wide
     variety of plant species. Exogenous application of BRs to plants at
     nanomolar to micromolar concentrations has a broad spectrum of growth
     responses, such as stem elongation, inhibition of root growth, promotion
     of cell division and enhancement of stress resistance, brought about by
     changes in enzyme activity and gene expression. In the last years,
     biochemical and genetic analysis provided compelling evidence for an
     essential role of BRs in plant development. In this paper, we review our
     synthetic methods to obtain BRs analogues and report the scope of
     antiviral activity of these compounds against RNA and DNA viruses.
     of the compounds showed selectivity indexes (SI) 10- to 18- fold higher
     than ribavirin, a broad spectrum antiviral compound, when tested against
     Junin virus (JV) (Arenaviridae); a good antiviral activity against measles
     virus (MV) (Paramixoviridae), with SI values also higher than ribavirin
     used as reference drug, and a similar or lower activity against herpes
     simplex type 1 and 2 (HSV-1 and HSV-2) (Herpesviridae) when compared to
     foscarnet or acyclovir, respectively. Structure activity relationship
     studies (SAR) are analyzed, in order to detect which stereochemistry, type
     and position of functional groups are needed to develop a selective class
     of virus inhibitors. . COPYRGT. 2004 Bentham Science Publishers Ltd.
     Medical Descriptors:
CT
     antiviral activity
     drug isolation
     plant
     stem elongation
     root growth
     cell division
     plant stress
     enzyme activity
     gene expression
```

```
chemical analysis
genetic analysis
plant development
RNA virus
DNA virus
drug selectivity
Junin virus
Arenavirus
Measles virus
Paramyxovirus
Herpes simplex virus 1
Herpes simplex virus 2
structure activity relation
stereochemistry
drug classification
drug synthesis
nonhuman
review
Drug Descriptors:
  *brassinosteroid: AN, drug analysis
  *brassinosteroid: DV, drug development
  *brassinosteroid: PD, pharmacology
lactone
ribavirin: CM, drug comparison
  ribavirin: PD, pharmacology
foscarnet: CM, drug comparison
  foscarnet: PD, pharmacology
aciclovir: CM, drug comparison
  aciclovir: PD, pharmacology
epoxide: AN, drug analysis
epoxide: DV, drug development
phytosterol: DV, drug development
  phytosterol: PD, pharmacology
orthoesterol A: DV, drug development
  orthoesterol A: PD, pharmacology
orthoesterol B: DV, drug development
  orthoesterol B: PD, pharmacology
orthoesterol C: DV, drug development
  orthoesterol C: PD, pharmacology
weibensterol A: DV, drug development
  weibensterol A: PD, pharmacology
weibensterol B: DV, drug development
  weibensterol B: PD, pharmacology
brassinolide: AN, drug analysis
brassinolide: CM, drug comparison
brassinolide: DV, drug development
 brassinolide: PD, pharmacology
stigmasterol: AN, drug analysis
stigmasterol: DV, drug development
2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha stigmastan 6 one: AN, drug
analysis
2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha stigmastan 6 one: CM, drug
comparison
2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha stigmastan 6 one: DV, drug
development
  2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha stigmastan 6 one: PD,
pharmacology
2 alpha 3 alpha 22,23 tetrahydroxy homo 7 oxastigmastan 6 one: AN, drug
analysis
2 alpha 3 alpha 22,23 tetrahydroxy homo 7 oxastigmastan 6 one: CM, drug
comparison
2 alpha 3 alpha 22,23 tetrahydroxy homo 7 oxastigmastan 6 one: DV, drug
development
  2 alpha 3 alpha 22,23 tetrahydroxy homo 7 oxastigmastan 6 one: PD,
pharmacology
2,3,22,23 bisacetyliden 2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha
```

```
sigmastan 6 one: AN, drug analysis
2,3,22,23 bisacetyliden 2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha
sigmastan 6 one: CM, drug comparison
2,3,22,23 bisacetyliden 2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha
sigmastan 6 one: DV, drug development
  2,3,22,23 bisacetyliden 2 alpha 3 alpha 22,23 tetrahydroxy 5 alpha
sigmastan 6 one: PD, pharmacology
2 alpha 3 alpha dihydroxy 5 alpha stigmast 22 en 6 one: AN, drug analysis
2 alpha 3 alpha dihydroxy 5 alpha stigmast 22 en 6 one: CM, drug
comparison
2 alpha 3 alpha dihydroxy 5 alpha stigmast 22 en 6 one: DV, drug
development
  2 alpha 3 alpha dihydroxy 5 alpha stigmast 22 en 6 one: PD,
pharmacology
2 alpha 3 alpha dihydroxy 6 alpha fluoro 5 alpha stigmast 22 ene: AN, drug
analysis
2 alpha 3 alpha dihydroxy 6 alpha fluoro 5 alpha stigmast 22 ene: CM, drug
comparison
2 alpha 3 alpha dihydroxy 6 alpha fluoro 5 alpha stigmast 22 ene: DV, drug
development
  2 alpha 3 alpha dihydroxy 6 alpha fluoro 5 alpha stigmast 22 ene: PD,
pharmacology
3 beta acetoxy 22,23 dihydroxy 5 alpha stigmastan 6 one: AN, drug analysis
3 beta acetoxy 22,23 dihydroxy 5 alpha stigmastan 6 one: CM, drug
comparison
3 beta acetoxy 22,23 dihydroxy 5 alpha stigmastan 6 one: DV, drug
development
  3 beta acetoxy 22,23 dihydroxy 5 alpha stigmastan 6 one: PD,
pharmacology
3 beta acetoxy 22,23 trihydroxy 5 alpha stigmastan 6 one: AN, drug
analysis
3 beta acetoxy 22,23 trihydroxy 5 alpha stigmastan 6 one: CM, drug
comparison
3 beta acetoxy 22,23 trihydroxy 5 alpha stigmastan 6 one: DV, drug
development
  3 beta acetoxy 22,23 trihydroxy 5 alpha stigmastan 6 one: PD,
pharmacology
3 alpha 22,23 trihydroxy 5 alpha stigmastan 6 one: CM, drug comparison
  3 alpha 22,23 trihydroxy 5 alpha stigmastan 6 one: PD,
pharmacology
3 alpha fluoro 22,23 dihydroxy 5 alpha stigmastan 6 one: AN, drug analysis
3 alpha fluoro 22,23 dihydroxy 5 alpha stigmastan 6 one: CM, drug
comparison
3 alpha fluoro 22,23 dihydroxy 5 alpha stigmastan 6 one: DV, drug
development
  3 alpha fluoro 22,23 dihydroxy 5 alpha stigmastan 6 one: PD,
pharmacology
2 alpha 3 alpha 5 alpha 22,23 pentahydroxystigmatan 6 one: AN, drug
analysis
2 alpha 3 alpha 5 alpha 22,23 pentahydroxystigmatan 6 one: CM, drug
comparison
2 alpha 3 alpha 5 alpha 22,23 pentahydroxystigmatan 6 one: DV, drug
development
  2 alpha 3 alpha 5 alpha 22,23 pentahydroxystigmatan 6 one: PD,
pharmacology
3 beta acetoxy 5 alpha 22,23 trihydrostigmastan 6 one: AN, drug analysis
3 beta acetoxy 5 alpha 22,23 trihydrostigmastan 6 one: CM, drug comparison
3 beta acetoxy 5 alpha 22,23 trihydrostigmastan 6 one: DV, drug
development
  3 beta acetoxy 5 alpha 22,23 trihydrostigmastan 6 one: PD,
pharmacology
3 beta bromo 5 alpha hydroxystigmast 22 en 6 one: AN, drug analysis
3 beta bromo 5 alpha hydroxystigmast 22 en 6 one: CM, drug comparison
3 beta bromo 5 alpha hydroxystigmast 22 en 6 one: DV, drug development
  3 beta bromo 5 alpha hydroxystigmast 22 en 6 one: PD, pharmacology
3 beta bromo 5 alpha 22,23 trihydroxystigmastan 6 one: AN, drug analysis
```

```
3 beta bromo 5 alpha 22,23 trihydroxystigmastan 6 one: CM, drug comparison
     3 beta bromo 5 alpha 22,23 trihydroxystigmastan 6 one: DV, drug
    development
       3 beta bromo 5 alpha 22,23 trihydroxystigmastan 6 one: PD,
     pharmacology
     3 beta fluoro 5 alpha 22,23 trihydroxystigmastan 6 one: AN, drug analysis
     3 beta fluoro 5 alpha 22,23 trihydroxystigmastan 6 one: CM, drug
     comparison
     3 beta fluoro 5 alpha 22,23 trihydroxystigmastan 6 one: DV, drug
     development
       3 beta fluoro 5 alpha 22,23 trihydroxystigmastan 6 one: PD,
     pharmacology
     3 beta 5 alpha dihydroxystigmast 22 en 6 one: AN, drug analysis
     3 beta 5 alpha dihydroxystigmast 22 en 6 one: CM, drug comparison
     3 beta 5 alpha dihydroxystigmast 22 en 6 one: DV, drug development
       3 beta 5 alpha dihydroxystigmast 22 en 6 one: PD, pharmacology
     unindexed drug
     Drug Descriptors:
CT
       unclassified drug
     (lactone) 1338-03-0; (ribavirin) 36791-04-5; (foscarnet) 4428-95-9;
RN
     (aciclovir) 59277-89-3; (brassinolide) 72962-43-7;
     (stigmasterol) 83-48-7
L55 ANSWER 3 OF 14 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.
     on STN
     2004032208 EMBASE
AN
     Steroid-hormone rapid actions, membrane receptors and a conformational
TI
     ensemble model.
     Norman A.W.; Mizwicki M.T.; Norman D.P.G.
ΑU
     A.W. Norman, Department of Biochemistry, University of California,
CS
     Riverside, CA 92521, United States. Anthony.norman@ucr.edu
     Nature Reviews Drug Discovery, (2004) Vol. 3, No. 1, pp. 27-41.
SO
     Refs: 118
     ISSN: 1474-1776 CODEN: NRDDAG
     United Kingdom
CY
     Journal; General Review
\mathtt{DT}
             Endocrinology
FS
     003
             Clinical Biochemistry
     029
            Pharmacology
     030
     037
             Drug Literature Index
     English
LA
     English
SL
     Entered STN: 20040129
ED
     Last Updated on STN: 20040129
     Steroid hormones can act as chemical messenger in a wide range od species
AB
     and target tissues to produce both slow genomic responses, and rapid
     non-genomic responses. Although it is clear that genomic responses to
     steroid hormones are mediated by the formation of a complex of the hormone
     and its cognate steroid-hormone nuclear receptor, new evidence indicates
     that rapid responses are mediated by a variety of receptor types
     associated with the plasma membrane or its caveolae components,
     potentially including a membrane-associated nuclear receptor. This review
     summarizes our current knowledge of membrane-associated steroid receptors,
     as well as details of structure-function relationships between steroid
     hormones and the ligand-binding domains of their nuclear and
     membrane-associated receptors. Furthermore, a new receptor conformational
     ensemble model is presented that suggests how the same receptor could
     produce both rapid and genomic responses. It is apparent that there is a
     cornucopia of new drug development opportunities in these areas.
     Medical Descriptors:
CT
     hormone action
     protein function
     conformational transition
     genomics
     complex formation
     cell membrane
```

```
caveola
    structure activity relation
    ligand binding
    protein domain
    tissue distribution
    drug targeting
    protein targeting
    drug receptor binding
    binding affinity
    human
    nonhuman
    human cell
    animal cell
    review
    priority journal
    Drug Descriptors:
    *steroid hormone: AN, drug analysis
      *steroid hormone: PD, pharmacology
    *membrane receptor: EC, endogenous compound
    *steroid receptor: EC, endogenous compound
    cell nucleus receptor: EC, endogenous compound
    estradiol: AN, drug analysis
      estradiol: PD, pharmacology
    androgen: AN, drug analysis
       androgen: PD, pharmacology
    alfacalcidol: AN, drug analysis
      alfacalcidol: PD, pharmacology
    glucocorticoid: AN, drug analysis
       glucocorticoid: PD, pharmacology
    mineralocorticoid: AN, drug analysis
      mineralocorticoid: PD, pharmacology
    thyroid hormone: AN, drug analysis
       thyroid hormone: PD, pharmacology
    peroxisome proliferator activated receptor: EC, endogenous compound
    retinoid: AN, drug analysis
       retinoid: PD, pharmacology
      brassinosteroid: AN, drug analysis
      brassinosteroid: PD, pharmacology
    testosterone: AN, drug analysis
       testosterone: PD, pharmacology
     progesterone: AN, drug analysis
       progesterone: PD, pharmacology
    hydrocortisone: AN, drug analysis
       hydrocortisone: PD, pharmacology
     aldosterone: AN, drug analysis
       aldosterone: PD, pharmacology
     retinoic acid: AN, drug analysis
       retinoic acid: PD, pharmacology
     liothyronine: AN, drug analysis
       liothyronine: PD, pharmacology
     ecdysone: AN, drug analysis
       ecdysone: PD, pharmacology
     brassinolide: AN, drug analysis
       brassinolide: PD, pharmacology
     ethinylestradiol: AN, drug analysis
       ethinylestradiol: PD, pharmacology
     vitamin D: AN, drug analysis
       vitamin D: PD, pharmacology
     (estradiol) 50-28-2; (alfacalcidol) 41294-56-8; (testosterone) 58-22-0;
     (progesterone) 57-83-0; (hydrocortisone) 50-23-7; (aldosterone) 52-39-1,
     6251-69-0; (retinoic acid) 302-79-4; (liothyronine) 6138-47-2, 6893-02-3;
     (brassinolide) 72962-43-7; (ethinylestradiol) 57-63-6
L55 ANSWER 4 OF 14 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.
     on STN
     2003369601 EMBASE
```

RN

AN

```
Structure-activity relationship studies in a set of new
TI
     brassinosteroid derivatives assayed against herpes simplex virus
     type 1 and 2 in cell cultures.
     Talarico L.B.; Ramirez J.A.; Galagovsky L.R.; Wachsman M.B.
AU
     Argentina. wachsman@qb.fcen.uba.ar
CS
     Medicinal Chemistry Research, (2002) Vol. 11, No. 8, pp. 434-444.
SO
     Refs: 20
     ISSN: 1054-2523 CODEN: MCREEB
     United States
CY
DT
     Journal; Article
             Pharmacology
FS
     030
             Drug Literature Index
     037
     English
LA
SL
     English
ED
     Entered STN: 20030925
     Last Updated on STN: 20030925
     Thirty-seven brassinosteroid derivatives were tested for their
AB
     antiviral activity against herpes simplex vires (HSV) type 1 and
     twenty-seven against HSV type 2, via a vires yield reduction assay.
     of the assayed compounds show selectivity indexes (SI) higher than those
     obtained with the reference drug, stigmasterol. The compounds that
     possessed a better structure-activity relationship are 6b
     [(22\dot{S}, 23S) - 3\beta-bromo-5\alpha, 22, 23-trihydroxystigmastan-6-one], 7b
     [(225,235)-3\beta, 5\alpha, 22,23-tetrahydroxystigmastan-6-one] and 12b
     [(22S, 23S) - 5\alpha - \text{fluor} - 3\beta, 22, 23-trihydroxy-stigmastan-6-one] with
     SI values of 100, 80 and 109 for HSV-1 and 71, 40 and 27 for HSV-2,
     respectively.
     Medical Descriptors:
CT
     *structure activity relation
     *antiviral activity
     Herpes simplex virus 1
     Herpes simplex virus 2
     cell culture
     drug selectivity
     nonhuman
     controlled study
     animal cell
     article
     Drug Descriptors:
       *brassinosteroid: AN, drug analysis
       *brassinosteroid: PD, pharmacology
     stigmasterol: AN, drug analysis
       stigmasterol: PD, pharmacology
     3 beta hydroxystigmasta 5,22 diene: AN, drug analysis
       3 beta hydroxystigmasta 5,22 diene: PD, pharmacology
     9 (2 hydroxyethoxymetyl) guanine: AN, drug analysis
       9 (2 hydroxyethoxymetyl) guanine: PD, pharmacology
     2alpha, 3alpha, 22, 23 tetrahydroxy 5alpha stigmastan 6 one: AN, drug
     analysis
       2alpha, 3alpha, 22, 23 tetrahydroxy 5alpha stigmastan 6 one: PD,
     pharmacology
     2alpha, 3alpha, 5alpha, 22, 23 pentahydroxystigmastan 6 one: AN, drug analysis
       2alpha, 3alpha, 5alpha, 22, 23 pentahydroxystigmastan 6 one: PD,
     pharmacology
     3beta acetoxy 22,23 dihydroxy 5alpha stigmastan 6 one: AN, drug analysis
       3beta acetoxy 22,23 dihydroxy 5alpha stigmastan 6 one: PD,
     pharmacology
     3beta acetoxy 5alpha, 22, 23 trihydroxystigmastan 6 one: AN, drug analysis
       3beta acetoxy 5alpha, 22, 23 trihydroxystigmastan 6 one: PD,
     pharmacology
     3beta bromo 22,23 diḥydroxy 5alpha stigmastan 6 one: AN, drug analysis
       3beta bromo 22,23 dihydroxy 5alpha stigmastan 6 one: PD,
     pharmacology
     3beta bromo 5alpha, 22, 23 trihydroxystigmastan 6 one: AN, drug analysis
       3beta bromo 5alpha, 22, 23 trihydroxystigmastan 6 one: PD,
     pharmacology
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3beta, 5alpha, 22, 23 tetrahydroxystigmastan 6 one: AN, drug analysis
       3beta, 5alpha, 22, 23 tetrahydroxystigmastan 6 one: PD, pharmacology
     3beta, 5alpha dihydroxystigmast 22 en 6 one: AN, drug analysis
       3beta,5alpha dihydroxystigmast 22 en 6 one: PD, pharmacology
     2alpha, 3alpha, 22, 23 tetrahydroxy beta homo 7 oxastigmastan 6 one: AN, drug
     analysis
       2alpha, 3alpha, 22, 23 tetrahydroxy beta homo 7 oxastigmastan 6 one: PD,
     pharmacology
     5alpha fluorostigmasta 2,22 dien 6 one: AN, drug analysis
       5alpha fluorostigmasta 2,22 dien 6 one: PD, pharmacology
     3beta fluoro 5alpha chlorostigmast 22 en 6 one: AN, drug analysis
       3beta fluoro 5alpha chlorostigmast 22 en 6 one: PD, pharmacology
     5alpha fluoro 3beta 22,23 trihydroxystigmastan 6 one: AN, drug analysis
       5alpha fluoro 3beta 22,23 trihydroxystigmastan 6 one: PD,
     pharmacology
     2alpha, 3alpha dihydroxystigmast 22 en 6 one: AN, drug analysis
       2alpha, 3alpha dihydroxystigmast 22 en 6 one: PD, pharmacology
     3beta bromo 5alpha hydroxystigmast 22 en 6 one: AN, drug analysis
       3beta bromo 5alpha hydroxystigmast 22 en 6 one: PD, pharmacology
     3beta fluoro 22,23 dihydroxystigmastan 6 one: AN, drug analysis
       3beta fluoro 22,23 dihydroxystigmastan 6 one: PD, pharmacology
     3beta fluoro 5alpha 22,23 dihydroxystigmastan 6 one: AN, drug analysis
       3beta fluoro 5alpha 22,23 dihydroxystigmastan 6 one: PD,
     pharmacology
     3alpha fluoro 22,23 dihydroxystigmastan 6 one: AN, drug analysis
       3alpha fluoro 22,23 dihydroxystigmastan 6 one: PD, pharmacology
     3beta bromo 5alpha chloro 6beta hydroxystigmast 22 ene: AN, drug analysis
       3beta bromo 5alpha chloro 6beta hydroxystigmast 22 ene: PD,
     pharmacology
     stigmasta 4,22 dien 3 one: AN, drug analysis
       stigmasta 4,22 dien 3 one: PD, pharmacology
     2alpa, 3alpha dihydroxy 6alpha fluoro 5alpha stigmast 22 ene: AN, drug
     analysis
       2alpa, 3alpha dihydroxy 6alpha fluoro 5alpha stigmast 22 ene: PD,
     pharmacology
     3alpha, 22, 23 trihydroxy 5alpha stigmastan 6 one: AN, drug analysis
       3alpha, 22, 23 trihydroxy 5alpha stigmastan 6 one: PD, pharmacology
     6beta hydroxy 5alpha stigmast 22 en 3 one: AN, drug analysis
       6beta hydroxy 5alpha stigmast 22 en 3 one: PD, pharmacology
     2alpha, 3alpha, 22, 23 tetrahydroxy 5alpha fluorostigmastan 6 one: AN, drug
     analysis
       2alpha, 3alpha, 22, 23 tetrahydroxy 5alpha fluorostigmastan 6 one: PD,
     pharmacology
     3beta, 22, 23 trihydroxy 5alpha stigmastan 6 one: AN, drug analysis
       3beta, 22, 23 trihydroxy 5alpha stigmastan 6 one: PD, pharmacology
     3alpha, 5alpha, 22, 23 tetrahydroxystigmastan 6 one: AN, drug analysis
       3alpha, 5alpha, 22, 23 tetrahydroxystigmastan 6 one: PD, pharmacology
     unindexed drug
     unclassified drug
     (stigmasterol) 83-48-7
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     M.B. Wachsman, Laboratorio de Virologia, Departamento de Quimica
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     wachsman@qb.fcen.uba.ar
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